Solid Waste



Best Proposed Demonstrated

Available Technology (BDAT) Background Document for K001

Volume 16

DRAFT REPORT

BEST DEMONSTRATED AVAILABLE TECHNOLOGY (BDAT) BACKGROUND DOCUMENT FOR K001

(WOOD PRESERVING INDUSTRY)

U.S. Environmental Protection Agency
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BDAT Background Document for K001

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EXECUTIVE SUMMARY

BDAT Treatment Standards for KOOl

Pursuant to the Hazardous and Solid Waste Amendments (HSWA) enacted on November 8, 1984, and in accordance with the procedures for establishing treatment standards under section 3004(m) of the Resource Conservation and Recovery Act (RCRA), the Environmental Protection Agency (EPA) is proposing treatment standards for the listed waste K001, based on the performance of treatment technologies determined by the Agency to represent Best Demonstrated Available Technology (BDAT). According to 40 CFR 261.32 waste code K001 is defined as "bottom sediment sludge from the treatment of wastewaters from wood preserving processes that use creosote and/or pentachlorophenol."

These BDAT treatment standards represent instantaneous maximum acceptable concentration levels for disposal of these wastes in units designated as land disposal units according to 40 CFR Part 268. Wastes that, as generated, contain the regulated constituents at concentrations which do not exceed the treatment standards are not restricted from land disposal units. The Agency has chosen to set levels for these wastes rather than designating the use of a specific treatment technology. The Agency believes that this approach allows the generators of these wastes a greater degree of flexibility in selecting a technology or train of technologies that can achieve these levels. The proposed effective date is August 8, 1988.

BDAT treatment standards for nonwastewater KOOl wastes are proposed based on performance data of rotary kiln incineration with stabilization of the nonwastewater residual. For the wastewater residual (i.e., scrubber water), the standard is based on the performance of a treatment train consisting of chemical precipitation and sludge filtration. Section 3 describes all applicable treatment technologies and presents treatment performance data which the Agency considered when developing the KOOl treatment standards.

Treatment standards have been proposed for a total of six organic constituents and three metals; the Agency believes that these constituents are indicators of effective treatment for all of the BDAT list constituents that have been identified as being present in K001 wastes. The BDAT list organic constituents that are proposed for regulation are naphthalene, pentachlorophenol, phenanthrene, pyrene, toluene, and xylenes (total). The BDAT list metals that are proposed for regulation are copper, lead, and zinc. The Agency has recently become aware of data showing that dioxins and furans may be present in some wood preserving wastes. EPA has not had an ample opportunity to evaluate these data. When the Agency completes its analysis of available data, it will consider the regulation of these constituents. A detailed discussion of the selection of regulated constituents is presented in Section 5 of this document.

The following tables list the specific BDAT treatment standards for K001 wastes. The Agency is proposing standards based on the analysis of total concentration for organic constituents and based on the analysis of TCLP extracts from K001 nonwastewaters. Standards are based on analysis of total concentration for K001 wastewaters. The units for total constituent concentration are in parts per million (mg/kg) on a weight by weight basis for nonwastewaters. The units for TCLP extract analysis are in parts per million (mg/l) on a weight by volume basis.

BDAT TREATMENT STANDARDS FOR KOOL

Nonwastewater

Regulated Constituent	Total Concentration (mg/kg)	TCLP Concentration (mg/l)
Naphtha lene	7 98	NA
Pentachlorophenol	36 75	NA .
Phenanthrene	7.98	NA
Pyrene	7.28	NA
Toluene	0.143	NA
Xylenes (total)	0 162	NA
Copper	NA	0.71
Lead	NA	0.53
Zinc	NA	0.086

NA = Not Applicable

Wastewater

Regulated Constituent	Total Concentration (mg/l)	TCLP Concentration (mg/1)
Naphthaìene	0 148	NA NA
Pentach lorophenol	0 875	NA
Pnenanthrene	0 148	NA NA
Pyrene	0 140	NA
To luene	0 143	NA
Xylenes (total)	0.161	NA
Copper	0 42	NA
Lead	0.037	NA
Zinc	1.0	NA

NA = Not applicable

1. INTRODUCTION

This section of the background document presents a summary of the legal authority pursuant to which the BDAT treatment standards were developed, a summary of EPA's promulgated methodology for developing BDAT, and finally a discussion of the petition process that should be followed to request a variance from the BDAT treatment standards.

1.1 Legal Background

1.1.1 Requirements Under HSWA

The Hazardous and Solid Waste Amendments of 1984 (HSWA), which were enacted on November 8, 1984, and which amended the Resource Conservation and Recovery Act of 1976 (RCRA), impose substantial new responsibilities on those who handle hazardous waste. In particular, the amendments require the Agency to promulgate regulations that restrict the land disposal of untreated hazardous wastes. In its enactment of HSWA, Congress stated explicitly that "reliance on land disposal should be minimized or eliminated, and land disposal, particularly landfill and surface impoundment, should be the least favored method for managing hazardous wastes" (RCRA section 1002(b)(7), 42 U.S.C. 6901(b)(7)).

One part of the amendments specifies dates on which particular groups of untreated hazardous wastes will be prohibited from land disposal unless "it has been demonstrated to the Administrator, to a reasonable degree of certainty, that there will be no migration of hazardous constituents from the disposal unit or injection zone for as long as the wastes remain hazardous" (RCRA section 3004(d)(1), (e)(1), (g)(5), 42 U.S.C. 6924(d)(1), (e)(1), (g)(5)).

For the purpose of the restrictions, HSWA defines land disposal "to include, but not be limited to, any placement of . . . hazardous waste in a landfill, surface impoundment, waste pile, injection well, land treatment facility, salt dome formation, salt bed formation, or underground mine or cave" (RCRA section 3004(k), 42 U.S.C. 6924(k)). Although HSWA defines land disposal to include injection wells, such disposal of solvents, dioxins, and certain other wastes, known as the California List wastes, is covered on a separate schedule (RCRA section 3004(f)(2), 42 U.S.C. 6924 (f)(2)). This schedule requires that EPA develop land disposal restrictions for deep well injection by August 8, 1988.

The amendments also require the Agency to set "levels or methods of treatment, if any, which substantially diminish the toxicity of the waste or substantially reduce the likelihood of migration of hazardous constituents from the waste so that short-term and long-term threats to human health and the environment are minimized" (RCRA section 3004(m)(1), 42 U.S.C. 6924 (m)(1)). Wastes that meet treatment standards established by EPA are not prohibited and may be land disposed. In setting treatment standards for listed or characteristic wastes, EPA may establish different standards for particular wastes within a single waste code with differing treatability characteristics. One such characteristic is the physical form of the waste. This frequently leads to different standards for wastewaters and nonwastewaters.

Alternatively, EPA can establish a treatment standard that is applicable to more than one waste code when, in EPA's judgment, all the waste can be treated to the same concentration. In those instances where a generator can demonstrate that the standard promulgated for the generator's waste cannot be achieved, the Agency also can grant a variance from a treatment standard by revising the treatment standard for that particular waste through rulemaking procedures. (A further discussion of treatment variances is provided in Section 1.3.)

The land disposal restrictions are effective when promulgated unless the Administrator grants a national variance and establishes a different date (not to exceed 2 years beyond the statutory deadline) based on "the earliest date on which adequate alternative treatment, recovery, or disposal capacity which protects human health and the environment will be available" (RCRA section 3004(h)(2), 42 U.S.C. 6924 (h)(2)).

If EPA fails to set a treatment standard by the statutory deadline for any hazardous waste in the First Third or Second Third of the schedule (see Section 1.1.2), the waste may not be disposed in a landfill or surface impoundment unless the facility is in compliance with the minimum technological requirements specified in section 3004(o) of RCRA. In addition, prior to disposal, the generator must certify to the Administrator that the availability of treatment capacity has been investigated, and it has been determined that disposal in a landfill or surface impoundment is the only practical alternative to treatment currently available to the generator. This restriction on the use of

landfills and surface impoundments applies until EPA sets a treatment standard for the waste or until May 8, 1990, whichever is sooner. If the Agency fails to set a treatment standard for any ranked hazardous waste by May 8, 1990, the waste is automatically prohibited from land disposal unless the waste is placed in a land disposal unit that is the subject of a successful "no migration" demonstration (RCRA section 3004(g), 42 U.S.C. 6924(g)). "No migration" demonstrations are based on case-specific petitions that show there will be no migration of hazardous constituents from the unit for as long as the waste remains hazardous.

1.1.2 Schedule for Developing Restrictions

Under section 3004(g) of RCRA, EPA was required to establish a schedule for developing treatment standards for all wastes that the Agency had listed as hazardous by November 8, 1984. Section 3004(g) required that this schedule consider the intrinsic hazards and volumes associated with each of these wastes. The statute required EPA to set treatment standards according to the following schedule:

- 1. Solvents and dioxins standards must be promulgated by November 8, 1986;
- 2. The "California List" must be promulgated by July 8, 1987;
- At least one-third of all listed hazardous wastes must be promulgated by August 8, 1988 (First Third);
- 4. At least two-thirds of all listed hazardous wastes must be promulgated by June 8, 1989 (Second Third); and
- 5. All remaining listed hazardous wastes and all hazardous wastes identified as of November 8, 1984, by one or more of the characteristics defined in 40 CFR Part 261 must be promulgated by May 8, 1990 (Third Third).

The statute specifically identified the solvent wastes as those covered under waste codes F001, F002, F003, F004, and F005; it identified the dioxin-containing hazardous wastes as those covered under waste codes F020, F021, F022, and F023.

Wastes collectively known as the California List wastes, defined under section 3004(d) of HSWA, are liquid hazardous wastes containing metals, free cyanides, PCBs, corrosives (i.e., a pH less than or equal to 2.0), and any liquid or nonliquid hazardous waste containing halogenated organic compounds (HOCs) above 0.1 percent by weight. Rules for the California List were proposed on December 11, 1986, and final rules for PCBs, corrosives, and HOC-containing wastes were established August 12, 1987. In that rule, EPA elected not to establish standards for metals. Therefore, the statutory limits became effective.

On May 28, 1986, EPA published a final rule (51 FR 19300) that delineated the specific waste codes that would be addressed by the First Third, Second Third, and Third Third. This schedule is incorporated into 40 CFR 268.10, 268.11, and 268.12.

1.2 <u>Summary of Promulgated BDAT Methodology</u>

In a November 7, 1986, rulemaking, EPA promulgated a technology-based approach to establishing treatment standards under section 3004(m). Section 3004(m) also specifies that treatment standards must "minimize" long- and short-term threats to human health and the environment arising from land disposal of hazardous wastes.

Congress indicated in the legislative history accompanying the HSWA that "[t]he requisite levels of [sic] methods of treatment established by the Agency should be the best that has been demonstrated to be achievable," noting that the intent is "to require utilization of available technology" and not a "process which contemplates technology-forcing standards" (Vol. 130 Cong. Rec. S9178 (daily ed., July 25, 1984)). EPA has interpreted this legislative history as suggesting that Congress considered the requirement under section 3004(m) to be met by application of the best demonstrated and achievable (i.e., available) technology prior to land disposal of wastes or treatment residuals. Accordingly, EPA's treatment standards are generally based on the performance of the best demonstrated available technology (BDAT) identified for treatment of the hazardous constituents. This approach involves the identification of potential treatment systems, the determination of whether they are demonstrated and available, and the collection of treatment data from well-designed and well-operated systems.

The treatment standards, according to the statute, can represent levels or methods of treatment, if any, that substantially diminish the toxicity of the waste or substantially reduce the likelihood of migration of hazardous constituents. Wherever possible, the Agency prefers to establish BDAT treatment standards as "levels" of treatment (i.e., performance standards), rather than adopting an approach that would require the use of specific treatment "methods." EPA believes that concentration-based treatment levels offer the regulated community greater

flexibility to develop and implement compliance strategies, as well as an incentive to develop innovative technologies.

1.2.1 Waste Treatability Group

In developing the treatment standards, EPA first characterizes the waste(s). As necessary, EPA may establish treatability groups for wastes having similar physical and chemical properties. That is, if EPA believes that wastes represented by different waste codes could be treated to similar concentrations using identical technologies, the Agency combines the codes into one treatability group. EPA generally considers wastes to be similar when they are both generated from the same industry and from similar processing stages. In addition, EPA may combine two or more separate wastes into the same treatability group when data are available showing that the waste characteristics affecting performance are similar or that one waste would be expected to be less difficult to treat.

Once the treatability groups have been established, EPA collects and analyzes data on identified technologies used to treat the wastes in each treatability group. The technologies evaluated must be demonstrated on the waste or a similar waste and must be available for use.

1.2.2 Demonstrated and Available Treatment Technologies

Consistent with legislative history, EPA considers demonstrated technologies to be those that are used to treat the waste of interest or a similar waste with regard to parameters that affect treatment selection (see November 7, 1986, 51 FR 40588). EPA also will consider as treatment those technologies used to separate or otherwise process chemicals and

other materials. Some of these technologies clearly are applicable to waste treatment, since the wastes are similar to raw materials processed in industrial applications.

For most of the waste treatability groups for which EPA will promulgate treatment standards, EPA will identify demonstrated technologies either through review of literature related to current waste treatment practices or on the basis of information provided by specific facilities currently treating the waste or similar wastes.

In cases where the Agency does not identify any facilities treating wastes represented by a particular waste treatability group, EPA may transfer a finding of demonstrated treatment. To do this, EPA will compare the parameters affecting treatment selection for the waste treatability group of interest to other wastes for which demonstrated technologies already have been determined. The parameters affecting treatment selection and their use for this waste are described in Section 3.2 of this document. If the parameters affecting treatment selection are similar, then the Agency will consider the treatment technology also to be demonstrated for the waste of interest. For example, EPA considers rotary kiln incineration to be a demonstrated technology for many waste codes containing hazardous organic constituents, high total organic content, and high filterable solids content, regardless of whether any facility is currently treating these wastes. The basis for this determination is data found in literature and data generated by EPA confirming the use of rotary kiln incineration on wastes having the above characteristics.

If no commercial treatment or recovery operations are identified for a waste or wastes with similar physical or chemical characteristics that affect treatment selection, the Agency will be unable to identify any demonstrated treatment technologies for the waste, and, accordingly, the waste will be prohibited from land disposal (unless handled in accordance with the exemption and variance provisions of the rule). The Agency is, however, committed to establishing treatment standards as soon as new or improved treatment processes are demonstrated (and available).

Operations only available at research facilities, pilot- and bench-scale operations, will not be considered in identifying demonstrated treatment technologies for a waste because these technologies would not necessarily be "demonstrated." Nevertheless, EPA may use data generated at research facilities in assessing the performance of demonstrated technologies.

As discussed earlier, Congress intended that technologies used to establish treatment standards under section 3004(m) be not only "demonstrated," but also available. To decide whether demonstrated technologies may be considered "available," the Agency determines whether they (1) are commercially available and (2) substantially diminish the toxicity of the waste or substantially reduce the likelihood of migration of hazardous constituents from the waste.

EPA will only set treatment standards based on a technology that meets the above criteria. Thus, the decision to classify a technology as "unavailable" will have a direct impact on the treatment standard. If

the best technology is unavailable, the treatment standard will be based on the next best treatment technology determined to be available. To the extent that the resulting treatment standards are less stringent, greater concentrations of hazardous constituents in the treatment residuals could be placed in land disposal units.

There also may be circumstances in which EPA concludes that for a given waste none of the demonstrated treatment technologies are "available" for purposes of establishing the 3004(m) treatment performance standards. Subsequently, these wastes will be prohibited from continued placement in or on the land unless managed in accordance with applicable exemptions and variance provisions. The Agency is, however, committed to establishing new treatment standards as soon as new or improved treatment processes become "available."

- (1) Proprietary or patented processes. If the demonstrated treatment technology is a proprietary or patented process that is not generally available, EPA will not consider the technology in its determination of the treatment standards. EPA will consider proprietary or patented processes available if it determines that the treatment method can be purchased or licensed from the proprietor or is a commercially available treatment. The services of the commercial facility offering this technology often can be purchased even if the technology itself cannot be purchased.
- (2) <u>Substantial treatment</u>. To be considered "available," a demonstrated treatment technology must "substantially diminish the

toxicity" of the waste or "substantially reduce the likelihood of migration of hazardous constituents" from the waste in accordance with section 3004(m). By requiring that substantial treatment be achieved in order to set a treatment standard, the statute ensures that all wastes are adequately treated before being placed in or on the land and ensures that the Agency does not require a treatment method that provides little or no environmental benefit. Treatment will always be deemed substantial if it results in nondetectable levels of the hazardous constituents of concern. If nondetectable levels are not achieved, then a determination of substantial treatment will be made on a case-by-case basis. This approach is necessary because of the difficulty of establishing a meaningful guideline that can be applied broadly to the many wastes and technologies to be considered. EPA will consider the following factors in an effort to evaluate whether a technology provides substantial treatment on a case-by-case basis:

- Number and types of constituents treated;
- Performance (concentration of the constituents in the treatment residuals); and
- Percent of constituents removed.

If none of the demonstrated treatment technologies achieve substantial treatment of a waste, the Agency cannot establish treatment standards for the constituents of concern in that waste.

1.2.3 Collection of Performance Data

Performance data on the demonstrated available technologies are evaluated by the Agency to determine whether the data are representative

of well-designed and well-operated treatment systems. Only data from well-designed and well-operated systems are included in determining BDAT. The data evaluation includes data already collected directly by EPA and/or data provided by industry. In those instances where additional data are needed to supplement existing information, EPA collects additional data through a sampling and analysis program. The principal elements of this data collection program are: (1) identification of facilities for site visits, (2) an engineering site visit, (3) a Sampling and Analysis Plan, (4) a sampling visit, and (5) an Onsite Engineering Report.

(1) <u>Identification of facilities for site visits</u>. To identify facilities that generate and/or treat the waste of concern, EPA uses a number of information sources. These include Stanford Research Institute's Directory of Chemical Producers; EPA's Hazardous Waste Data Management System (HWDMS); the 1986 Treatment, Storage, Disposal Facility (TSDF) National Screening Survey; and EPA's Industry Studies Data Base. In addition, EPA contacts trade associations to inform them that the Agency is considering visits to facilities in their industry and to solicit their assistance in identifying facilities for EPA to consider in its treatment sampling program.

After identifying facilities that treat the waste, EPA uses this hierarchy to select sites for engineering visits: (1) generators treating single wastes on site; (2) generators treating multiple wastes together on site; (3) commercial treatment, storage, and disposal facilities

(TSDFs); and (4) EPA in-house treatment. This hierarchy is based on two concepts: (1) to the extent possible, EPA should develop treatment standards from data produced by treatment facilities handling only a single waste, and (2) facilities that routinely treat a specific waste have had the best opportunity to optimize design parameters. Although excellent treatment can occur at many facilities that are not high in this hierarchy, EPA has adopted this approach to avoid, when possible, ambiguities related to the mixing of wastes before and during treatment.

When possible, the Agency will evaluate treatment technologies using commercially operated systems. If performance data from properly designed and operated commercial treatment methods for a particular waste or a waste judged to be similar are not available, EPA may use data from research facilities operations. Whenever research facility data are used, EPA will explain in the preamble and background document why such data were used and will request comments on the use of such data.

Although EPA's data bases provide information on treatment for individual wastes, the data bases rarely provide data that support the selection of one facility for sampling over another. In cases where several treatment sites appear to fall into the same level of the hierarchy, EPA selects sites for visits strictly on the basis of which facility could most expeditiously be visited and later sampled if justified by the engineering visit.

(2) Engineering site visit. Once a treatment facility has been selected, an engineering site visit is made to confirm that a candidate for sampling meets EPA's criteria for a well-designed facility and to ensure that the necessary sampling points can be accessed to determine operating parameters and treatment effectiveness. During the visit, EPA also confirms that the facility appears to be well operated, although the actual operation of the treatment system during sampling is the basis for EPA's decisions regarding proper operation of the treatment unit. In general, the Agency considers a well-designed facility to be one that contains the unit operations necessary to treat the various hazardous constituents of the waste, as well as to control other nonhazardous materials in the waste that may affect treatment performance.

In addition to ensuring that a system is reasonably well designed, the engineering visit examines whether the facility has a way to measure the operating parameters that affect performance of the treatment system during the waste treatment period. For example, EPA may choose not to sample a treatment system that operates in a continuous mode, for which an important operating parameter cannot be continuously recorded. In such systems, instrumentation is important in determining whether the treatment system is operating at design values during the waste treatment period.

(3) <u>Sampling and Analysis Plan</u>. If after the engineering site visit the Agency decides to sample a particular plant, the Agency will then develop a site-specific Sampling and Analysis Plan (SAP) according to the Generic Quality Assurance Project Plan for the Land Disposal Restriction

Program ("BDAT"), EPA/530-SW-87-011. In brief, the SAP discusses where the Agency plans to sample, how the samples will be taken, the frequency of sampling, the constituents to be analyzed and the method of analysis, operational parameters to be obtained, and specific laboratory quality control checks on the analytical results.

The Agency will generally produce a draft of the site-specific Sampling and Analysis Plan within 2 to 3 weeks of the engineering visit. The draft of the SAP is then sent to the plant for review and comment. With few exceptions, the draft SAP should be a confirmation of data collection activities discussed with the plant personnel during the engineering site visit. EPA encourages plant personnel to recommend any modifications to the SAP that they believe will improve the quality of the data.

It is important to note that sampling of a plant by EPA does not mean that the data will be used in the development of treatment standards for BDAT. EPA's final decision on whether to use data from a sampled plant depends on the actual analysis of the waste being treated and on the operating conditions at the time of sampling. Although EPA would not plan to sample a facility that was not ostensibly well designed and well operated, there is no way to ensure that at the time of the sampling the facility will not experience operating problems. Additionally, EPA statistically compares its test data to suitable industry-provided data, where available, in its determination of what data to use in developing treatment standards. The methodology for comparing data is presented later in this section.

(Note: Facilities wishing to submit data for consideration in the development of BDAT standards should, to the extent possible, provide sampling information similar to that acquired by EPA. Such facilities should review the Generic Quality Assurance Project Plan for the Land Disposal Restriction Program ("BDAT"), which delineates all of the quality control and quality assurance measures associated with sampling and analysis. (Quality assurance and quality control procedures are summarized in Section 1.2.6 of this document.)

(4) <u>Sampling visit</u>. The purpose of the sampling visit is to collect samples that characterize the performance of the treatment system and to document the operating conditions that existed during the waste treatment period. At a minimum, the Agency attempts to collect sufficient samples of the untreated waste and solid and liquid treatment residuals so that variability in the treatment process can be accounted for in the development of the treatment standards. To the extent practicable, and within safety constraints, EPA or its contractors collect all samples and ensure that chain-of-custody procedures are conducted so that the integrity of the data is maintained.

In general, the samples collected during the sampling visit will have already been specified in the SAP. In some instances, however, EPA will not be able to collect all planned samples because of changes in the facility operation or plant upsets; EPA will explain any such deviations from the SAP in its follow-up Onsite Engineering Report.

(5) Onsite Engineering Report. EPA summarizes all its data collection activities and associated analytical results for testing at a facility in a report referred to as the Onsite Engineering Report (OER). This report characterizes the waste(s) treated, the treated residual concentrations, the design and operating data, and all analytical results including methods used and accuracy results. This report also describes any deviations from EPA's suggested analytical methods for hazardous wastes (see Test Methods for Evaluating Solid Waste, SW-846, Third Edition, November 1986).

After the Onsite Engineering Report is completed, the report is submitted to the plant for review. This review provides the plant with a final opportunity to claim any information contained in the report as confidential. Following the review and incorporation of comments, as appropriate, the report is made available to the public with the exception of any material claimed as confidential by the plant.

- 1.2.4 Hazardous Constituents Considered and Selected for Regulation
- (1) <u>Development of BDAT list</u>. The list of hazardous constituents within the waste codes that are targeted for treatment is referred to by the Agency as the BDAT constituent list. This list, provided as Table 1-1, is derived from the constituents presented in 40 CFR Part 261, Appendices VII and VIII, as well as several ignitable constituents used as the basis of listing wastes as F003 and F005. These sources provide a comprehensive list of hazardous constituents specifically regulated under RCRA. The BDAT list consists of those constituents that can be analyzed using methods published in SW-846, Third Edition.

Table 1-1 BDAT Constituent List

BDAT	D	0.1.0
reference	Parameter	CAS no.
no		
	<u>Volatiles</u>	
222	Acetone	67-64-1
1.	Acetonitrile	75-05-8
2.	Acrolein	107-02-8
3.	Acrylonitrile	107-13-1
4.	Benzene	71-43-2
5.	Bromodichloromethane	75-27 - 4
6.	Bromomethane	74-83-9
223	n-Butyl alcohol	71-36-3
7.	Carbon tetrachloride	56-23-5
8.	Carbon disulfide	75-15-0
9.	Chlorobenzene	108-90-7
10.	2-Chloro-1.3-butadiene	126-99-8
11	Chlorodibromomethane	124-48-1
12	Chloroethane	75-00-3
13.	2-Chloroethyl vinyl ether	110-75-8
14.	Chloroform	67-66-3
15	Chloromethane	74-87-3
16.	3-Chloropropene	107-05-1
17.	1,2-Dibromo-3-chloropropane	96-12-8
18.	1.2-Dibromoethane	106-93-4
19.	Dibromomethane	74-95-3
20	Trans-1,4-Dichloro-2-butene	110-57-6
21.	Dichlorodifluoromethane	75-71-8
22.	1,1-Dichloroethane	75-34-3
23.	1,2-Dichloroethane	107-06-2
24.	1,1-Dichloroethylene	75-35-4
25.	Trans-1,2-Dichloroethene	156-60-5
26.	1,2-Dichloropropane	78-87-5
27.	Trans-1,3-Dichloropropene	10061-02-6
28.	cis-1,3-Dichloropropene	10061-01-5
29.	1,4-Dioxane	123-91-1
224.	2-Ethoxyethanol	110-80-5
225.	Ethyl acetate	141-78-6
226.	Ethyl benzene	100-41-4
30.	Ethyl cyanide	107-12-0
227.	Ethyl ether	60-29-7
31.	Ethyl methacrylate	97-63-2
214	Ethylene oxide	75-21-8
32.	Iodomethane	74-88-4

Table 1-1 (continued)

BDAT	****	
reference	Parameter	CAS no.
no	Taraneter	CAS IIU.

	<u>Volatiles</u> (continued)	
33.	Isobutyl alcohol	78-83-1
228.	Methano l	67-56-1
34.	Methyl ethyl ketone	78-93-3
229.	Methyl isobutyl ketone	108-10-1
35.	Methyl methacrylate	80-62-6
37.	Methacrylonitrile	126-98-7
38.	Methylene chloride	75-09-2
230.	2-Nitropropane	79-46-9
39.	Pyridine	110-86-1
40.	1,1,1,2-Tetrachloroethane	630-20-6
41.	1,1,2,2-Tetrachloroethane	79-34-6
42.	Tetrachloroethene	127-18-4
43	Toluene	108-88-3
44.	Tribromomethane	75-25-2
45	1,1,1-Trichloroethane	71-55-6
46.	1,1,2-Trichloroethane	79-00-5
47	Trichloroethene	79-01-6
48.	Trichloromonofluoromethane	75-69-4
49.	1,2,3-Trichloropropane	96-18-4
231.	1,1,2-Trichloro-1,2,2-trifluoro- ethane	76-13-1
50.	Vinyl chloride	75-01-4
215	1,2-Xylene	97-47-6
216.	1,3-Xylene	108-38-3
217.	1,4-Xylene	106-44-5
	<u>Semivolatiles</u>	
51.	Acenaphthalene	208-96-8
52	Acenaphthene	83-32-9
53.	Acetophenone	96-86-2
54.	2-Acetylaminofluorene	53-96-3
55.	4-Aminobiphenyl	92-67-1
56.	Aniline	62-53-3
57.	Anthracene	120-12-7
58.	Aramite	140-57-8
59	Benz(a)anthracene	56-55-3
218	Benzal chloride	98-87-3
60.	Benzeneth101	108-98-5
61.	De leted	
62.	Benzo(a)pyrene	50-32-8

Table 1-1 (continued)

BDAT reference no	Parameter	CAS no.
		-
	<u>Semivolatiles</u> (continued)	
63	Benzo(b)fluoranthene	205-99-2
64	Benzo(ghi)perylene	191-24-2
65.	Benzo(k)fluoranthene	207-08-9
66	p-Benzoquinone	106-51-4
67.	Bis(2-chloroethoxy)methane	111-91-1
68.	Bis(2-chloroethyl)ether	111-44-4
69.	Bis(2-chloroisopropyl)ether	39638-32-9
70.	Bis(2-ethylhexyl)phthalate	117-81-7
71	4-Bromophenyl phenyl ether	101-55-3
72.	Butyl benzyl phthalate	85-68-7
73.	2-sec-Butyl-4,6-dinitrophenol	88-85-7
74.	p-Chloroaniline	106-47-8
75	Chlorobenzilate	510-15-6
76	p-Chloro-m-cresol	59-50-7
77	2-Chloronaphthalene	91-58-7
78	2-Chlorophenol	95-57-8
79	3-Chloropropionitrile	542-76-7
80.	Chrysene	218-01-9
81	ortho-Creso	95-48-7
82	para-Cresol	106-44-5
232	Cyclohexanone	108-94-1
83	Dibenz(a,h)anthracene	53-70-3
84	Dibenzo(a,e)pyrene	192-65-4
85	Dibenzo(a,i)pyrene	189-55-9
86	m-Dichlorobenzene	541-73-1
87	o-Dichlorobenzene	95-50-1
88	p-Dichlorobenzene	106-46-7
89	3,3'-Dichloropenzidine	91-94-1
90	2,4-Dichlorophenol	120-83-2
91	2,6-Dichlorophenol	87-65-0
92.	Diethyl phthalate	84-66-2
93	3,3'-Dimethoxybenzidine	119-90-4
94	p-Dimethylaminoazobenzene	60-11-7
95	3,3'~Dimethylbenzidine	119-93-7
96	2,4-Dimethylphenol	105-67-9
97.	Dimethyl phthalate	131-11-3
98	Di-n-butyl pnthalate	84-74-2
99	1.4-Dinitropenzene	100-25-4
100.	4,6-Dinitro-o-cresol	534-52-1
100.	2,4-Dinitrophenol	51-28-5

Table 1-1 (continued)

BDAT		
reference	Parameter	CAS no.
no		
	Somewolatalos (acatemical)	
	Semivolatiles (continued)	
102	2,4-Dinitrotoluene	121-14-2
103	2,6-Dinitrotoluene	606-20-2
104	Di-n-octyl phthalate	117-84-0
105.	Di-n-propylnitrosamine	621-64-7
106	Diphenylamine	122-39-4
219.	Diphenylnitrosamine	86-30-6
107.	1,2-Diphenylhydrazine	122-66-7
108	Fluoranthene	206-44-0
109	Fluorene	86-73-7
110	Hexachlorobenzene	118-74-1
111	Hexachlorobutadiene	87- 68-3
112	Hexachlorocyclopentadiene	77-4 7-4
113	Hexachloroethane	67-72-1
114	Hexachlorophene	70-30-4
115	Hexachloropropene	1888-71-7
116	Indeno(1,2,3-cd)pyrene	193-39-5
117	Isosafrole	120-58-1
118	Methapyr:lene	91-80-5
119	3-Methylcholanthrene	56-49-5
120	4,4'-Methylenebis	
	(2-chloroaniline)	101-14-4
36.	Methyl methanesulfonate	66-27-3
121	Naphtha lene	91-20-3
122.	1,4-Naphthoquinone	130-15-4
123	1-Naphthylamine	134-32-7
124	2-Naphthylamine	91-59-8
125.	p-Nitroaniline	100-01-6
126.	Nitrobenzene	98-95-3
127.	4-Nitrophenol	100-02-7
128	N-Nitrosodi-n-butylamine	924-16-3
129.	N-Nitrosodiethylamine	55-18-5
130	N-Nitrosodimethylamine	62-75-9
131.	N-Nitrosomethylethylamine	10595-95-6
132.	N-Nitrosomorpholine	59-89-2
133.	N-Nitrosopiperidine	100-75-4
134	n-Nitrosopyrrolidine	930-55-2
135	5-Nitro-o-toluidine	99-65-8
136	Pentachlorobenzene	608-93-5
137	Pentach loroethane	76-01-7
138	Pentachloronitrobenzene	82-68-8

Table 1-1 (continued)

BDAT reference	Parameter	CAS no
no.		
	<u>Semivolatiles</u> (continued)	
139.	Pentachlorophenol	87-86-5
140	Phenacet in	62-44-2
141	Phenanthrene	85-01-8
142	Pheno 1	108-95-2
220.	Phthalic anhydride	85-44-9
143.	2-Picoline	109-06-8
144.	Pronamide	23950-58-5
145.	Pyrene	129-00-0
146.	Resorcinol	108-46-3
147.	Safrole	94-59-7
148.	1,2,4,5-Tetrachlorobenzene	95-94-3
149.	2,3,4,6-Tetrachlorophenol	58-90-2
150.	1,2,4-Trichlorobenzene	120-82-1
151.	2,4,5-Trichlorophenol	95-95-4
152	2,4,6-Trichlorophenol	88-06-2
153.	Tris(2,3-dibromopropyl)	
	⁻ phosphate	126-72-7
	<u>Metals</u>	
154.	Antimony	7440-36-0
155.	Arsenic	7440-38-2
156.	Barıum	7440-39-3
157.	Beryllium	7440-41-7
158.	Cadmıum	7440-43-9
159.	Chromium (total)	7440-47-32
221.	Chromium (hexavalent)	-
160.	Copper	7440-50-8
161.	Lead	7439-92-1
162.	Mercury	7439-97-6
163.	Nickel	7440-02-0
164.	Selenium Salaaa	7782-49-2
165.	Silver	7440-22-4
166.	Thallium	7440-28-0
167.	Vanadium	7440-62-2
168.	Zinc	7440-66-6
	<u>Inorganics</u> other than metals	
169.	Cyanide	57-12-5
170.	Fluoride	16964-48-8
171.	Sulfide	8496-25-8

Table 1-1 (continued)

BDAT reference no	Parameter	CAS no.
110		
	Organochlorine pesticides	
172.	Aldrın	309-00-2
173.	a lpha-BHC	319-84-6
174.	beta-BHC	319-85-7
175.	delta-BHC	319-86-8
176.	gamma-BHC	58-89-9
177.	Chlordane	57 -74 -9
178.	DDD	72-54-8
179.	DDE	72-55-9
180.	DDT	50-29-3
181.	Dieldrin	60-57-1
182.	Endosulfan I	939-98-8
183.	Endosulfan II	33213-6-5
184.	Endrin	72-20-8
185.	Endrin aldehyde	7421-93-4
186	Heptachlor	76-44-8
187.	Heptachlor epoxide	1024-57-3
188.	Isodrin	465-73 -6
189.	Kepone	143-50-0
190. ,	Methoxyclor	72-43-5
191.	Toxaphene	8001-35-2
	Phenoxyacetic acid herbicides	
192.	2,4-Dichlorophenoxyacetic acid	94-75-7
193.	Silvex	93-72-1
194.	2,4,5-T	93-76-5
	Organophosphorous insecticides	
195	Disulfoton	298-04-4
196	Famphur	52-85-7
197.	Methyl parathion	298-00-0
198.	Parathion	56-38-2
199.	Phorate	298-02-2
	PCBs	
200.	Aroclor 1016	12674-11-2
201.	Aroclor 1221	11104-28-2
202.	Aroclor 1232	11141-16-5

Table 1-1 (continued)

BDAT reference no	Parameter	CAS no.
	PCBs (continued)	
203.	Aroclor 1242	53469-21-9
204.	Araclor 1248	12672-29-6
205.	Aroclor 1254	11097-69-1
206.	Aroclor 1260	11096-82-5
	Dioxins and furans	
207.	Hexachlorodibenzo-p-dioxins	-
208	Hexachlorodibenzofurans	-
209.	Pentachlorodibenzo-p-dioxins	-
210.	Pentachlorodibenzofurans	-
211.	Tetrachlorodibenzo-p-dioxins	-
212	Tetrachlorodibenzofurans	-
213.	2,3,7,8-Tetrachlorodibenzo-p-dioxin	1746-01-6

The initial BDAT constituent list was published in EPA's Generic Quality Assurance Project Plan, March 1987 (EPA/530-SW-87-011).

Additional constituents will be added to the BDAT constituent list as more key constituents are identified for specific waste codes or as new analytical methods are developed for hazardous constituents. For example, since the list was published in March 1987, 18 additional constituents (hexavalent chromium, xylenes (all three isomers), benzal chloride, phthalic anhydride, ethylene oxide, acetone, n-butyl alcohol, 2-ethoxyethanol, ethyl acetate, ethyl benzene, ethyl ether, methanol, methyl isobutyl ketone, 2-nitropropane, 1,1,2-trichloro-1,2,2-trifluoroethane, and cyclohexanone) have been added to the list.

Chemicals are listed in Appendix VIII if they are shown in scientific studies to have toxic, carcinogenic, mutagenic, or teratogenic effects on humans or other life-forms, and they include such substances as those identified by the Agency's Carcinogen Assessment Group as being carcinogenic. Including a constituent in Appendix VIII means that the constituent can be cited as a basis for listing toxic wastes.

Although Appendix VII, Appendix VIII, and the F003 and F005 ignitables provide a comprehensive list of RCRA-regulated hazardous constituents, not all of the constituents can be analyzed in a complex waste matrix. Therefore, constituents that could not be readily analyzed in an unknown waste matrix were not included on the initial BDAT list. As mentioned above, however, the BDAT constituent list is a continuously growing list that does not preclude the addition of new constituents when analytical methods are developed.

There are five major reasons that constituents were not included on the BDAT constituent list:

- 1. Constituents are unstable. Based on their chemical structure, some constituents will either decompose in water or will ionize. For example, maleic anhydride will form maleic acid when it comes in contact with water and copper cyanide will ionize to form copper and cyanide ions. However, EPA may choose to regulate the decomposition or ionization products.
- 2. EPA-approved or verified analytical methods are not available. Many constituents, such as 1,3,5-trinitrobenzene, are not measured adequately or even detected using any of EPA's analytical methods published in SW-846 Third Edition.
- 3. The constituent is a member of a chemical group designated in Appendix VIII as not otherwise specified (N.O.S.). Constituents listed as N.O.S., such as chlorinated phenols, are a generic group of some types of chemicals for which a single analytical procedure is not available. The individual members of each such group need to be listed to determine whether the constituents can be analyzed. For each N.O.S. group, all those constituents that can be readily analyzed are included in the BDAT constituent list.
- 4. Available analytical procedures are not appropriate for a complex waste matrix. Some compounds, such as auramine, can be analyzed as a pure constituent. However, in the presence of other constituents, the recommended analytical method does not positively identify the constituent. The use of high pressure liquid chromatography (HPLC) presupposes a high expectation of finding the specific constituents of interest. In using this procedure to screen samples, protocols would have to be developed on a case-specific basis to verify the identity of constituents present in the samples. Therefore, HPLC is not an appropriate analytical procedure for complex samples containing unknown constituents.
- 5. Standards for analytical instrument calibration are not commercially available. For several constituents, such as benz(c)acridine, commercially available standards of a "reasonably" pure grade are not available. The unavailability of a standard was determined by a review of catalogs from specialty chemical manufacturers.

Two constituents (fluoride and sulfide) are not specifically included in Appendices VII and VIII; however, these compounds are included on the BDAT list as indicator constituents for compounds from Appendices VII and VIII such as hydrogen fluoride and hydrogen sulfide, which ionize in water.

The BDAT constituent list presented in Table 1-1 is divided into the following nine groups:

- Volatile organics;
- Semivolatile organics;
- Metals:
- Other inorganics;
- Organochlorine pesticides;
- Phenoxyacetic acid herbicides;
- Organophosphorous insecticides;
- PCBs; and
- Dioxins and furans.

The constituents were placed in these categories based on their chemical properties. The constituents in each group are expected to behave similarly during treatment and are also analyzed, with the exception of the metals and inorganics, by using the same analytical methods.

(2) <u>Constituent selection analysis</u>. The constituents that the Agency selects for regulation in each treatability group are, in general, those found in the untreated wastes at treatable concentrations. For certain waste codes, the target list for the untreated waste may have been shortened (relative to analyses performed to test treatment technologies) because of the extreme unlikelihood that the constituent will be present.

In selecting constituents for regulation, the first step is to summarize all the constituents that were found in the untreated waste at treatable concentrations. This process involves the use of the statistical analysis of variance (ANOVA) test, described in Section 1.2.6, to determine if constituent reductions were significant. The Agency interprets a significant reduction in concentration as evidence that the technology actually "treats" the waste.

There are some instances where EPA may regulate constituents that are not found in the untreated waste but are detected in the treated residual. This is generally the case where presence of the constituents in the untreated waste interferes with the quantification of the constituent of concern. In such instances, the detection levels of the constituent are relatively high, resulting in a finding of "not detected" when, in fact, the constituent is present in the waste.

After determining which of the constituents in the untreated waste are present at treatable concentrations, EPA develops a list of potential constituents for regulation. The Agency then reviews this list to determine if any of these constituents can be excluded from regulation because they would be controlled by regulation of other constituents in the list.

EPA performs this indicator analysis for two reasons: (1) it reduces the analytical cost burdens on the treater and (2) it facilitates implementation of the compliance and enforcement program. EPA's rationale for selection of regulated constituents for this waste code is presented in Section 5 of this background document.

(3) Calculation of standards. The final step in the calculation of the BDAT treatment standard is the multiplication of the average treatment value by a factor referred to by the Agency as the variability factor. This calculation takes into account that even well-designed and well-operated treatment systems will experience some fluctuations in performance. EPA expects that fluctuations will result from inherent mechanical limitations in treatment control systems, collection of treated samples, and analysis of these samples. All of the above fluctuations can be expected to occur at well-designed and well-operated treatment facilities. Therefore, setting treatment standards utilizing a variability factor should be viewed not as a relaxing of section 3004(m) requirements, but rather as a function of the normal variability of the treatment processes. A treatment facility will have to be designed to meet the mean achievable treatment performance level to ensure that the performance levels remain within the limits of the treatment standard.

The Agency calculates a variability factor for each constituent of concern within a waste treatability group using the statistical calculation presented in Appendix A. The equation for calculating the variability factor is the same as that used by EPA for the development of numerous regulations in the Effluent Guidelines Program under the Clean Water Act. The variability factor establishes the instantaneous maximum based on the 99th percentile value.

There is an additional step in the calculation of the treatment standards in those instances where the ANOVA analysis shows that more

than one technology achieves a level of performance that represents BDAT. In such instances, the BDAT treatment standard is calculated by first averaging the mean performance value for each technology for each constituent of concern and then multiplying that value by the highest variability factor among the technologies considered. This procedure ensures that all the BDAT technologies used as the basis for the standards will achieve full compliance.

1.2.5 Compliance with Performance Standards

All the treatment standards reflect performance achieved by the best demonstrated available technology (BDAT). As such, compliance with these standards requires only that the treatment level be achieved prior to land disposal. It does not require the use of any particular treatment technology. While dilution of the waste as a means to comply with the standard is prohibited, wastes that are generated in such a way as to naturally meet the standard can be land disposed without treatment. With the exception of treatment standards that prohibit land disposal, all treatment standards proposed are expressed as a concentration level.

EPA has used both total constituent concentration and TCLP analyses of the treated waste as a measure of technology performance. EPA's rationale for when each of these analytical tests is used is explained in the following discussion.

For all organic constituents, EPA is basing the treatment standards on the total constituent concentration found in the treated waste. EPA based its decision on the fact that technologies exist to destroy the

various organics compounds. Accordingly, the best measure of performance would be the extent to which the various organic compounds have been destroyed or the total amount of constituent remaining after treatment. (NOTE: EPA's land disposal restrictions for solvent waste codes F001-F005 (51 FR 40572) use the TCLP value as a measure of performance. At the time that EPA promulgated the treatment standards for F001-F005, useful data were not available on total constituent concentrations in treated residuals and, as a result, the TCLP data were considered to be the best measure of performance.)

For all metal constituents, EPA is using both total constituent concentration and/or the TCLP as the basis for treatment standards. The total constituent concentration is being used when the technology basis includes a metal recovery operation. The underlying principle of metal recovery is the reduction of the amount of metal in a waste by separating the metal for recovery; therefore, total constituent concentration in the treated residual is an important measure of performance for this technology. Additionally, EPA also believes that it is important that any remaining metal in a treated residual waste not be in a state that is easily leachable; accordingly, EPA is also using the TCLP as a measure of performance. It is important to note that for wastes for which treatment standards are based on a metal recovery process, the facility has to comply with both the total constituent concentration and the TCLP prior to land disposal.

In cases where treatment standards for metals are not based on recovery techniques but rather on stabilization, EPA is using only the TCLP as a measure of performance. The Agency's rationale is that stabilization is not meant to reduce the concentration of metal in a waste but only to chemically minimize the ability of the metal to leach.

1.2.6 Identification of BDAT

- (1) <u>Screening of treatment data</u>. This section explains how the Agency determines which of the treatment technologies represent treatment by BDAT. The first activity is to screen the treatment performance data from each of the demonstrated and available technologies according to the following criteria:
 - Design and operating data associated with the treatment data must reflect a well-designed, well-operated system for each treatment data point. (The specific design and operating parameters for each demonstrated technology for this waste code are discussed in Section 3.2 of this document.)
 - 2. Sufficient QA/QC data must be available to determine the true values of the data from the treated waste. This screening criterion involves adjustment of treated data to take into account that the type value may be different from the measured value. This discrepancy generally is caused by other constituents in the waste that can mask results or otherwise interfere with the analysis of the constituent of concern.
 - 3. The measure of performance must be consistent with EPA's approach to evaluating treatment by type of constituents (e.g., total concentration data for organics, and total concentration and TCLP for metals in the leachate from the residual).

In the absence of data needed to perform the screening analysis, EPA will make decisions on a case-by-case basis as to whether to include the data. The factors included in this case-by-case analysis will be the

actual treatment levels achieved, the availability of the treatment data and their completeness (with respect to the above criteria), and EPA's assessment of whether the untreated waste represents the waste code of concern. EPA's application of these screening criteria for this waste code is provided in Section 4 of this background document.

(2) <u>Comparison of treatment data</u>. In cases in which EPA has treatment data from more than one technology following the screening activity, EPA uses the statistical method known as analysis of variance (ANOVA) to determine if one technology performs significantly better than the others. This statistical method (summarized in Appendix A) provides a measure of the differences between two data sets. If EPA finds that one technology performs significantly better (i.e., the data sets are not homogeneous), BDAT treatment standards are the level of performance achieved by the best technology multiplied by the corresponding variability factor for each regulated constituent.

If the differences in the data sets are not statistically significant, the data sets are said to be homogeneous. Specifically, EPA uses the analysis of variance to determine whether BDAT represents a level of performance achieved by only one technology or represents a level of performance achieved by more than one (or all) of the technologies. If the Agency finds that the levels of performance for one or more technologies are not statistically different, EPA averages the performance values achieved by each technology and then multiplies this value by the largest variability factor associated with any of the

acceptable technologies. A detailed discussion of the treatment selection method and an example of how EPA chooses BDAT from multiple treatment systems is provided in Section A-1.

(3) Quality assurance/quality control. This section presents the principal quality assurance/quality control (QA/QC) procedures employed in screening and adjusting the data to be used in the calculation of treatment standards. Additional QA/QC procedures used in collecting and screening data for the BDAT program are presented in EPA's Generic Quality Assurance Project Plan for Land Disposal Restrictions Program ("BDAT") (EPA/530-SW-87-011, March 1987).

To calculate the treatment standards for the Land Disposal Restriction Rules, it is first necessary to determine the recovery value for each constituent (the amount of constituent recovered after spiking, which is the addition of a known amount of the constituent, minus the initial concentration in the samples divided by the amount added) for a spike of the treated residual. Once the recovery value is determined, the following procedures are used to select the appropriate percent recovery value to adjust the analytical data:

If duplicate spike recovery values are available for the constituent of interest, the data are adjusted by the lowest available percent recovery value (i.e., the value that will yield the most conservative estimate of treatment achieved). However, if a spike recovery value of less than 20 percent is reported for a specific constituent, the data are not used to set treatment standards because the Agency does not have sufficient confidence in the reported value to set a national standard.

- 2. If data are not available for a specific constituent but are available for an isomer, then the spike recovery data are transferred from the isomer and the data are adjusted using the percent recovery selected according to the procedure described in (1) above.
- 3. If data are not available for a specific constituent but are available for a similar class of constituents (e.g., volatile organics, acid-extractable semivolatiles), then spike recovery data available for this class of constituents are transferred. All spike recovery values greater than or equal to 20 percent for a spiked sample are averaged and the constituent concentration is adjusted by the average recovery value. If spiked recovery data are available for more than one sample, the average is calculated for each sample and the data are adjusted by the lowest average value.
- 4. If matrix spike recovery data are not available for a set of data to be used to calculate treatment standards, then matrix spike recovery data are transferred from a waste that the Agency believes is a similar matrix (e.g., if the data are for an ash from incineration, then data from other incinerator ashes could be used). While EPA recognizes that transfer of matrix spike recovery data from a similar waste is not an exact analysis, this is considered the best approach for adjusting the data to account for the fact that most analyses do not result in extraction of 100 percent of the constituent. In assessing the recovery data to be transferred, the procedures outlined in (1), (2), and (3) above are followed.

The analytical procedures employed to generate the data used to calculate the treatment standards are listed in Appendix B of this document. In cases where alternatives or equivalent procedures and/or equipment are allowed in EPA's SW-846, Third Edition (November 1986) methods, the specific procedures and equipment used are also documented in this Appendix. In addition, any deviations from the SW-846, Third Edition, methods used to analyze the specific waste matrices are documented. It is important to note that the Agency will use the methods and procedures delineated in Appendix B to enforce the treatment

standards presented in Section 6 of this document. Accordingly, facilities should use these procedures in assessing the performance of their treatment systems.

- 1.2.7 BDAT Treatment Standards for "Derived-From" and "Mixed" Wastes
- (1) Wastes from treatment trains generating multiple residues. In a number of instances, the proposed BDAT consists of a series of operations, each of which generates a waste residue. For example, the proposed BDAT for a certain waste code is based on solvent extraction, steam stripping, and activated carbon adsorption. Each of these treatment steps generates a waste requiring treatment—a solvent—containing stream from solvent extraction, a stripper overhead, and spent activated carbon. Treatment of these wastes may generate further residues; for instance, spent activated carbon (if not regenerated) could be incinerated, generating an ash and possibly a scrubber water waste. Ultimately, additional wastes are generated that may require land disposal. With respect to these wastes, the Agency wishes to emphasize the following points:
 - 1. All of the residues from treating the original listed wastes are likewise considered to be the listed waste by virtue of the derived-from rule contained in 40 CFR Part 261.3(c)(2). (This point is discussed more fully in (2) below.) Consequently, all of the wastes generated in the course of treatment would be prohibited from land disposal unless they satisfy the treatment standard or meet one of the exceptions to the prohibition.
 - The Agency's proposed treatment standards generally contain a concentration level for wastewaters and a concentration level for nonwastewaters. The treatment standards apply to all of the wastes generated in treating the original prohibited waste. Thus, all solids generated from treating these wastes would have

to meet the treatment standard for nonwastewaters. All derived-from wastes meeting the Agency definition of wastewater (less than 1 percent TOC and less than 1 percent total filterable solids) would have to meet the treatment standard for wastewaters. EPA wishes to make clear that this approach is not meant to allow partial treatment in order to comply with the applicable standard.

- 3. The Agency has not performed tests, in all cases, on every waste that can result from every part of the treatment train. However, the Agency's treatment standards are based on treatment of the most concentrated form of the waste. Consequently, the Agency believes that the less concentrated wastes generated in the course of treatment will also be able to be treated to meet this value.
- (2) <u>Mixtures and other derived-from residues</u>. There is a further question as to the applicability of the BDAT treatment standards to residues generated not from treating the waste (as discussed above), but from other types of management. Examples are contaminated soil or leachate that is derived from managing the waste. In these cases, the mixture is still deemed to be the listed waste, either because of the derived-from rule (40 CFR Part 261.3(c)(2)(i)) or the mixture rule (40 CFR Part 261.3(a)(2)(iii) and (iv)) or because the listed waste is contained in the matrix (see, for example, 40 CFR Part 261.33(d)). The prohibition for the particular listed waste consequently applies to this type of waste.

The Agency believes that the majority of these types of residues can meet the treatment standards for the underlying listed wastes (with the possible exception of contaminated soil and debris for which the Agency is currently investigating whether it is appropriate to establish a separate treatability subcategorization). For the most part, these

residues will be less concentrated than the original listed waste. The Agency's treatment standards also make a generous allowance for process variability by assuming that all treatability values used to establish the standard are lognormally distributed. The waste also might be amenable to a relatively nonvariable form of treatment technology such as incineration. Finally, and perhaps most important, the rules contain a treatability variance that allows a petitioner to demonstrate that its waste cannot be treated to the level specified in the rule (40 CFR Part 268.44(a)). This provision provides a safety valve that allows persons with unusual waste matrices to demonstrate the appropriateness of a different standard. The Agency, to date, has not received any petitions under this provision (for example, for residues contaminated with a prohibited solvent waste), indicating, in the Agency's view, that the existing standards are generally achievable.

wastes. The Agency has been asked if and when residues from managing hazardous wastes, such as leachate and contaminated ground water, become subject to the land disposal prohibitions. Although the Agency believes this question to be settled by existing rules and interpretative statements, to avoid any possible confusion the Agency will address the question again.

Residues from managing First Third wastes, listed California List wastes, and spent solvent and dioxin wastes are all considered to be subject to the prohibitions for the underlying hazardous waste. Residues

from managing California List wastes likewise are subject to the California List prohibitions when the residues themselves exhibit a characteristic of hazardous waste. This determination stems directly from the derived-from rule in 40 CFR Part 261.3(c)(2) or, in some cases, from the fact that the waste is mixed with or otherwise contains the listed waste. The underlying principle stated in all of these provisions is that listed wastes remain listed until delisted.

The Agency's historic practice in processing delisting petitions that address mixing residuals has been to consider them to be the listed waste and to require that delisting petitioners address all constituents for which the derived-from waste (or other mixed waste) was listed. The language in 40 CFR Part 260.22(b) states that mixtures or derived-from residues can be delisted provided a delisting petitioner makes a demonstration identical to that which a delisting petitioner would make for the underlying waste. Consequently, these residues are treated as the underlying listed waste for delisting purposes. The statute likewise takes this position, indicating that soil and debris that are contaminated with listed spent solvents or dioxin wastes are subject to the prohibition for these wastes even though these wastes are not the originally generated waste, but rather are a residual from management (RCRA section 3004(e)(3)). It is EPA's view that all such residues are covered by the existing prohibitions and treatment standards for the listed hazardous waste that these residues contain and from which they are derived.

1.2.8 Transfer of Treatment Standards

EPA is proposing some treatment standards that are not based on testing of the treatment technology of the specific waste subject to the treatment standard. Instead, the Agency has determined that the constituents present in the subject waste can be treated to the same performance levels as those observed in other wastes for which EPA has previously developed treatment data. EPA believes that transferring treatment performance for use in establishing treatment standards for untested wastes is technically valid in cases where the untested wastes are generated from similar industries, have similar processing steps, or have similar waste characteristics affecting performance and treatment selection. Transfer of treatment standards to similar wastes or wastes from similar processing steps requires little formal analysis. However, in a case where only the industry is similar, EPA more closely examines the waste characteristics prior to deciding whether the untested waste constituents can be treated to levels associated with tested wastes.

EPA undertakes a two-step analysis when determining whether wastes generated by different processes within a single industry can be treated to the same level of performance. First, EPA reviews the available waste characteristic data to identify those parameters that are expected to affect treatment selection. EPA has identified some of the most important constituents and other parameters needed to select the treatment technology appropriate for a given waste. A detailed discussion of each analysis, including how each parameter was selected for each waste, can be found in Section 5 of this document.

Second, when an individual analysis suggests that an untested waste can be treated with the same technology as a waste for which treatment performance data are already available, EPA analyzes a more detailed list of constituents that represent some of the most important waste characteristics that the Agency believes will affect the performance of the technology. By examining and comparing these characteristics, the Agency determines whether the untested wastes will achieve the same level of treatment as the tested waste. Where the Agency determines that the untested waste is easier to treat than the tested waste, the treatment standards can be transferred. A detailed discussion of this transfer process for each waste can be found in later sections of this document.

1.3 Variance from the BDAT Treatment Standard

The Agency recognizes that there may exist unique wastes that cannot be treated to the level specified as the treatment standard. In such a case, a generator or owner/operator may submit a petition to the Administrator requesting a variance from the treatment standard. A particular waste may be significantly different from the wastes considered in establishing treatability groups because the waste contains a more complex matrix that makes it more difficult to treat. For example, complex mixtures may be formed when a restricted waste is mixed with other waste streams by spills or other forms of inadvertent mixing. As a result, the treatability of the restricted waste may be altered such that it cannot meet the applicable treatment standard.

Variance petitions must demonstrate that the treatment standard established for a given waste cannot be met. This demonstration can be

made by showing that attempts to treat the waste by available technologies were not successful or by performing appropriate analyses of the waste, including waste characteristics affecting performance, which demonstrate that the waste cannot be treated to the specified levels. Variances will not be granted based solely on a showing that adequate BDAT treatment capacity is unavailable. (Such demonstrations can be made according to the provisions in Part 268.5 of RCRA for case-by-case extensions of the effective date.) The Agency will consider granting generic petitions provided that representative data are submitted to support a variance for each facility covered by the petition.

Petitioners should submit at least one copy to:

The Administrator U.S. Environmental Protection Agency 401 M Street, S.W. Washington, DC 20460

An additional copy marked "Treatability Variance" should be submitted to:

Chief, Waste Treatment Branch Office of Solid Waste (WH-565) U.S. Environmental Protection Agency 401 M Street, S.W. Washington, DC 20460

Petitions containing confidential information should be sent with only the inner envelope marked "Treatability Variance" and "Confidential Business Information" and with the contents marked in accordance with the requirements of 40 CFR Part 2 (41 FR 36902, September 1, 1976, amended by 43 FR 4000).

The petition should contain the following information:

- 1. The petitioner's name and address.
- 2. A statement of the petitioner's interest in the proposed action.
- 3. The name, address, and EPA identification number of the facility generating the waste, and the name and telephone number of the plant contact.
- 4. The process(es) and feed materials generating the waste and an assessment of whether such process(es) or feed materials may produce a waste that is not covered by the demonstration.
- 5. A description of the waste sufficient for comparison with the waste considered by the Agency in developing BDAT, and an estimate of the average and maximum monthly and annual quantities of waste covered by the demonstration. (Note: The petitioner should consult the appropriate BDAT background document for determining the characteristics of the wastes considered in developing treatment standards.)
- 6. If the waste has been treated, a description of the system used for treating the waste, including the process design and operating conditions. The petition should include the reasons the treatment standards are not achievable and/or why the petitioner believes the standards are based on inappropriate technology for treating the waste. (Note: The petitioner should refer to the BDAT background document as guidance for determining the design and operating parameters that the Agency used in developing treatment standards.)
- 7. A description of the alternative treatment systems examined by the petitioner (if any); a description of the treatment system deemed appropriate by the petitioner for the waste in question; and, as appropriate, the concentrations in the treatment residual or extract of the treatment residual (i.e., using the TCLP, where appropriate, for stabilized metals) that can be achieved by applying such treatment to the waste.
- 8. A description of those parameters affecting treatment selection and waste characteristics that affect performance, including results of all analyses. (See Section 3.0 for a discussion of waste characteristics affecting performance that the Agency has identified for the technology representing BDAT.)
- 9. The dates of the sampling and testing. •
- 10. A description of the methodologies and equipment used to obtain representative samples.

- 11. A description of the sample handling and preparation techniques, including techniques used for extraction, containerization, and preservation of the samples.
- 12. A description of analytical procedures used, including QA/QC methods.

After receiving a petition for a variance, the Administrator may request any additional information or waste samples that may be required to evaluate and process the petition. Additionally, all petitioners must certify that the information provided to the Agency is accurate under 40 CFR Part 268.4(b).

In determining whether a variance will be granted, the Agency will first look at the design and operation of the treatment system being used. If EPA determines that the technology and operation are consistent with BDAT, the Agency will evaluate the waste to determine if the waste matrix and/or physical parameters are such that the BDAT treatment standards reflect treatment of this waste. Essentially, this latter analysis will concern the parameters affecting treatment selection and waste characteristics affecting performance parameters.

In cases where BDAT is based on more than one technology, the petitioner will need to demonstrate that the treatment standard cannot be met using any of the technologies, or that none of the technologies are appropriate for treatment of the waste. After the Agency has made a determination on the petition, the Agency's findings will be published in the <u>Federal Register</u>, followed by a 30-day period for public comment.

After review of the public comments, EPA will publish its final determination in the <u>Federal Register</u> as an amendment to the treatment standards in 40 CFR Part 268, Subpart D.

2. INDUSTRIES AFFECTED AND WASTE CHARACTERIZATION

The previous section discussed the BDAT program and the methodology used by the Agency to develop treatment standards. The purpose of this section is to describe the industry affected by the land disposal restrictions for K001 waste, the process generating the waste, and the available waste characterization data.

According to 40 CFR Part 261.32 (hazardous wastes from specific sources), the waste identified as K001 is specifically generated from the treatment of wastewaters from wood preserving processes that use creosote and/or pentachlorophenol. For the purpose of BDAT determination, the Agency has determined that K001 wastes generated from the use of creosote or pentachlorophenol based wood preservatives are similar and represent one treatability group.

2.1 <u>Industry Affected and Process Description</u>

The listed waste KOOl is generated in the wood preserving industry. The four digit Standard Industrial Classification (SIC) code most often reported for the wood preserving industry is 2491. The Agency estimates that at least 400 facilities have wood preserving processes that could potentially generate KOOl waste. Table 2-1 lists the number of facilities by State. Table 2-2 summarizes the number of facilities for each EPA Region. Figure 2-1 illustrates these data geographically on a map of the United States.

The preservation of wood using creosote and/or pentachlorophenol generates wastewaters containing hazardous constituents present in the preservatives. This process is illustrated in Figure 2-2. Creosote is a

Table 2-1 Number of Wood Preserving Facilities by State*

State	Facilities	State	Facilities	
AL (IV)	29	MT (VIII)		
AK (X)		NE (VII)		
AZ (IX)		NV (IX)		
AR (VI)	17	NH (I)		
CA (IX)	31	NJ (II)		
CO (VIII)		NM (VI)		
CT (I)		NY (II)		
DE (III)		NC(IV)	29	
DC (III)		ND (VIII)		
FL (IV)	25	OH (V)	11	
GA (IV)	30	OK (VI)		
HI (IX)		OR (X)	12	
ID (X)		PA (III)	16	
IL (V)	11	RI (1)		
IN (V)	13	SC (IV)	11	
IA (VII)		SD (VIII)		
KS (VII)		TN (IV)	7	
KY (IV)	12	TX (VI)	33	
LA (VI)	21	UT (VIII)		
ME (I)		VT (I)		
MD (III)		VA (III)	17	
MA (I)		WA (X)	22	
MI (V)		WV (III)	8	
MN (V)	12	WI (V)	11	
MS (IV)	27	WY (VIII)		
MO (VII)	22			

^{*} Includes data for SIC code 2491 only. Reference. 1982 Census of Manufactures

Table 2-2 Number of Wood Preserving Facilities by EPA Region*

EPA Region	Totals			
I	_			
11	-			
111	41			
IV	170			
٧	58			
VI	81			
A11	22			
VIII	-			
IX	31			
X	_34			
	437			

^{*}Includes data for SIC code 2491 only. Reference. 1982 Census of Manufactures.

FIGURE 2-1. WOOD PRESERVING FACILITIES BY STATE AND EPA REGION

DC

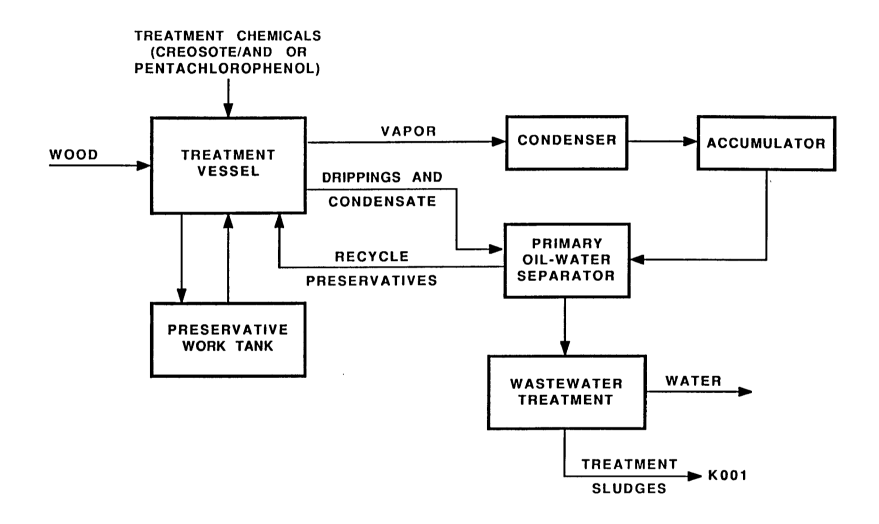


FIGURE 2-2. WOOD PRESERVING PROCESS

derivative of coal containing a wide range of constituents including, cresols, phenol, 2,4-dimethylphenol, naphthalene, benz(a) anthracene, benzo(a)pyrene, fluoranthene, benzo(b)fluoranthene, chrysene, benzo(a,h)anthracene, indeno(1,2,3-cd)pyrene, and acenaphthalene. The treatment by any means (including simple settling) of these wastewaters generates the listed waste K001.

The wood preserving process consists of two steps: (1) pretreatment of the wood to reduce its natural moisture content, and (2) impregnation of the wood with preservatives including creosote and/or pentachlorophenol. These agents are added to wood to increase its resistance to natural decay, attack by insects, and microorganisms. Drippings and condensed vapors generated during preservation treatment are sent to the oil-water separator. In the oil-water separator, wood treatment chemicals are recovered and recycled back to the preserving process. The wastewater, contaminated with components of creosote, pentachlorophenol, and/or other related compounds, is pumped to wastewater treatment. The treatment residual generated is the listed waste K001.

2.2 Waste Characterization

This section includes all waste characterization data available to the Agency for KOO1. An estimate of the major constituents that comprise the waste and their approximate concentrations is presented in Table 2-3. The percent concentration of each major constituent in the waste was determined by best estimates based on chemical analyses of KOO1 wastes from wood preservation using creosote and from pentachlorophenol

Table 2-3 Major Constituent Composition*
Untreated K001 Waste

Major Constituents	Concentration (%)			
Soil	35			
BDAT List Organic Constituents				
Naphthalene	4 0			
Phenanthrene	3.5			
Fluoranthene	2 0			
Acenaphthene	2.0			
Pyrene	1.5			
Fluorene	1.5			
Anthracene	1 0			
Pentachlorophenol	<1.0			
0tners	8.5			
Water	20			
Other Organic Compounds	14			
Wood Chips	5			
BDAT List Metals	_<1_			
	100%			

^{*} Percent concentrations presented here were determined from engineering judgment based on chemical analyses.

based treatment chemicals. The Agency has obtained compositional data from its own testing program and from numerous literature sources. The ranges of BDAT list constituents present in the wastes and other available data are presented in Table 2-4.

These data indicate the various BDAT list organic constituents present in K001 waste. These data display the wide ranges of concentrations of hazardous organics that may be present in the wastes. Such variations may be attributed to the type of preservative chemicals used and the type of wastewater treatment systems used. Generally, these K001 wastes contain numerous polynuclear aromatic compounds and chlorinated phenolics present in the wood preservatives. No characterization data identified in the literature for K001 had values for BDAT list metals. The BDAT list metals detected in samples of K001 waste collected by the Agency are presented in Section 3 of this document.

Analyses for dioxins and furans were performed by the Agency on K001 wastes collected for developing treatment standards. These compounds were not detected in any of the nine samples analyzed. The Agency has recently become aware of waste characterization data for wood preserving wastes showing that dioxins and furans may be present. The Agency is currently evaluating these data.

Table 2-4 BDAT List Constituent Composition

	Untreated K001 Waste Total Composition (ppm)						
BDAT Constituent Source of Data	(a)	(b)	(c)	(d)	(e)	(f)	(g)
Acenaphthene	-	3,000	-	-		15,000-21,000	13,000-18,000
Anthracene	8,410	-	-	-	-	7,300-15,000	8,500-13,000
Chrysene	-	45	9.29	4 5	2.1	4,100-4,800	<2,500-3,400
Fluoranthene	5,090	1,400	-	-	-	BDL	13,000-21,000
Naphtha lene	43,640	1,200	-	-	-	29,000-43.000	26,000-43,000
Pyrene	604	52	-	-	-	12,000-17,000	9,200-15,000
Phenanthrene	8,410	3,200	-	-	-	28,000-42,000	28,000-43,000
Pentachlorophenol	1 84	-	4 8	302	58	BDL	920-3,000
2,ă-dichlorophenol	1,650	-	-	-	-	BDL	BDL
p-chloro-m-cresol	1,690	-	-	-	-	BDL	BDL
2,4-dimethyl phenol	-	8.2	-	4 4	3.4	BDL	BDL
Benzo[g,h,ı] perylene	-	84	-	-	-	BDL	BDL
Fluorene	-	1,400	-	-	-	12,000-18,000	8,200-12,000
Dibez[a,h] anthracene	-	-	0.052	-	-	BDL	BDL
Benz[a] anthracene	-	-	1.25	3 7	0.149	BDL	<2,500-3,400
Benzo[a] pyrene	-	-	5 98	-	-	BDL	<250-340
Pheno 1	-	-	4.5	9 0	16	2,400-3,900	BDL
2-Chlorophenol	-	-	0 30	39	1.2	BDL	BDL
2,4,6-trichlorophenol	-	-	-	-	25	BDL	BDL
Benzo(b and/or k) fluoranthrene	-	-	-	_	-	BDL	940-2,300

^{- =} No Data

BDL = Below Detection Limit

⁽a) Reference - RCRA Background Listing Document for K001

⁽b) Reference - RCRA Background Listing Document for K001

⁽c) Reference - Myers 1979

⁽d) Reference - Myers 1979

⁽e) Reference - Myers 1979

⁽f) Reference - Onsite Engineering Report for K001-Creosote

⁽g) Reference - Onsite Engineering Report for K001-Pentachlorophenol

Table 2-4 BDAT List Constituent Composition (continued)

		Untreated KOOl Waste Total Composition (ppm)						
BDAT Constituent Source of Da	Data (h)	(1)	(1)	(k)	(1)	(m)	(n)	(0)
Acenaphthene	~	~	-	-	-	-	-	-
Anthracene	~	-	-	-	-	-	-	-
Chrysene	~	-	-	1-170		-	-	-
Fluoranthene	-	-	-	-	_	-	-	-
Naphtha lene	20,000	-	-	-	1.7-150	-	-	-
yrene	~	-	-	-	0.17-440	-	-	~
Phenanthrene	~	-	-	3.5-900	-	-	-	=
Pentachlorophenol 50	0,000-200,000	10,000	0 034	-	-	55-1,500	0 18-30	20,000-50,00
2,4-dichlorophenol	-	-	-	-	-	165	_	-
o-chloro-m-cresol	-	-	-	-	-	0 17	_	-
2.4-dimethyl phenol	-		-	~	-	-	-	-
Benzo[g,h,ı] perylene	-	-	-	-	-	-	-	-
luorene	-	-	-	0.93-560	~	-	-	-
Dibez[a,h] anthracene	~	-	-	-	-	-	-	-
Benz[a] anthracene	-	-	-	0.014-0 37	1-260	-	-	-
Benzo[a] pyrene	-	-	-	-	-	-	-	-
Phenol	-	10,000	5 043	-	-	<5.0	0.45-1 6	~
?-Chlorophenol	-	-	-	-	-	-	0 12-39 6	-
2,4,6-trichlorophenol	-	-	-	-	-	2.5	-	-
2,4-Dinitrophenol	-	-	0.024	-	~	-	0 3-0.8	-
"Creosote"	-	-	10	-	-	_	-	-

^{- =} No Data

BDL = Below Detection Limit

⁽h) Reference - USEPA 1980

⁽i) Reference ~ DPRA 1984

⁽j) Reference - Myers 1979

⁽k) Reference - K.W. Brown 1981

⁽¹⁾ Reference - Acurex 1982

⁽m) Reference - Acurex 1982

⁽n) Reference - Mitre 1981

⁽o) Reference - Illinois EPA 1983

3. APPLICABLE/DEMONSTRATED TREATMENT TECHNOLOGIES

In the previous section there was a discussion of the industries generating K001 and the untreated waste composition. This section describes the applicable treatment technologies, demonstrated treatment technologies, and available performance data for K001. The Agency identified applicable treatment technologies based on available waste composition data, contacts with industry, and technical publications. The technologies considered to be applicable to the untreated waste are those that treat hazardous organic compounds by reducing their concentration. Additionally, treatment residuals (wastewater and nonwastewater) are expected to contain metals in treatable concentrations. Treatment technologies applicable to reducing the concentration and/or leachability of metals in these residuals were also identified. Included in this section are discussions of those treatment technologies that have been demonstrated on a commercial basis. Treatment performance data collected by the Agency for these demonstrated technologies are also presented.

3.1 Applicable Treatment Technologies

The chemical composition of KOOl waste most directly affects the technologies applicable to the waste. As shown in Section 2, the waste primarily contains high concentrations of BDAT list organic constituents, high filterable solids content, moderate water content, and BDAT list metals at concentrations below one percent. Treatment technologies are needed for treatment of both BDAT list organics and BDAT list metals.

For BDAT list organics, the Agency has identified treatment technologies that may be applicable to KOOl because the technologies are designed to treat organic constituents in high filterable solids matrices. The technologies applicable to KOOl are those that destroy or remove the organics present in the untreated waste.

The Agency has identified the following treatment technologies as being applicable to BDAT list organic constituents in K001: incineration and fuel substitution. Incineration is a destruction technology that destroys organic constituents in wastes. Fuel substitution, like incineration, destroys the organic constituents of a waste while deriving a fuel value from the waste.

The goal of incineration is to thermally destroy (oxidize) the organic constituents of a waste. Typically, the types of incineration systems that are demonstrated on wastes include fluidized bed, rotary kiln, fixed hearth, and liquid injection. The Agency believes that the performance of rotary kiln incineration adequately represents the performance achievable by other thermal destruction technologies (including fuel substitution) that are well designed and well operated, and can handle sludges of this type. Rotary kiln incineration systems are designed specifically to handle sludges, solids, tarry wastes, and containerized liquids that are difficult to atomize through a liquid injector. Many rotary kiln incinerators are also designed to simultaneously incinerate other liquid wastes or supplemental fuel.

The Agency believes that solvent extraction may be applicable to K001 waste; however, EPA has not identified any facilities using solvent extraction on K001 or a similar waste. The Agency does not currently have sufficient information on waste parameters that affect treatment selection for solvent extraction to suggest that this technology is applicable for wastes similar to K001, accordingly, EPA does not consider solvent extraction to be an applicable technology.

Incineration technologies generally result in the formation of two treatment residuals: ash and scrubber water. For the BDAT list metals present in the wastewater residual (i.e., scrubber water), the applicable treatment technologies are chemical precipitation and filtration. Chemical precipitation removes dissolved metals from solution, and filtration removes suspended solids that result from the use of an underdesigned clarifier or from the generation of precipitates that do not settle easily. The filter cake generated from filtration contains BDAT list metals and requires stabilization before land disposal.

For the BDAT list metals present in the nonwastewater residuals (wastewater treatment filter cake and ash), the applicable treatment technologies are high temperature metals recovery and stabilization. High temperature metals recovery provides for recovery of metals from wastes primarily by volatilization of metals and subsequent condensation and collection steps. The process yields a metal product for reuse and reduces the amount of waste that requires land disposal. Stabilization

is designed to chemically and physically bind metal constituents of the waste into the microstructure of a cementitious matrix and thereby reduce their leaching potential. A variety of reagents, including Portland cements, cement kiln dust, hydrated limes, quick lime, fly ash, and other pozzalanic materials, have been demonstrated to act as binding reagents for various types of wastes containing metals.

3.2 <u>Demonstrated Treatment Technologies</u>

Of the two applicable technologies for BDAT list organic constituents, the Agency believes that rotary kiln incineration is demonstrated to treat KOOl since it is being used to treat wastes similar to KOOl with regard to parameters affecting treatment selection, including high concentrations of BDAT list organics and high filterable solids content. Fuel substitution is also demonstrated on KOOl because it is demonstrated on similar wastes with regard to parameters affecting treatment selection. Performance data for rotary kiln incineration of KOOl are presented in Tables 3-1 to 3-9, in Section 3.3 following the demonstrated treatment technology descriptions.

For KOOl wastewaters, the Agency believes that chemical precipitation and filtration are demonstrated because they are demonstrated on similar wastewater streams containing BDAT list metals. Treatment performance data for BDAT list metal constituents in wastewater treatment residuals from incineration of KOOl are presented in Table 3-14 in Section 3.3, following the descriptions of the demonstrated treatment technologies.

For BDAT list metals treatment, EPA believes that stabilization is demonstrated to treat K001 nonwastewater treatment residuals because it is being used to treat similar wastes with regard to parameters affecting treatment selection. The Agency does not believe that high temperature metals recovery is demonstrated on K001 waste residuals because it is not demonstrated specifically on K001 residuals or on similar wastes.

Specifically, at this time, EPA does not have treatment performance data for high temperature metals recovery of K001 filter cake, incinerator ash, or other wastes having similar types and concentrations of BDAT list metal constituents. Treatment performance data for stabilization of filter cake or incinerator ash from incineration of K001 are presented in Tables 3-15 through 3-17.

3.2.1 Incineration

This section addresses the commonly used incineration technologies: Liquid injection, rotary kiln, fluidized bed incineration, and fixed hearth. A discussion is provided regarding the applicability of these technologies, the underlying principles of operation, a technology description, waste characteristics that affect performance, and finally important design and operating parameters. As appropriate the subsections are divided by type of incineration unit.

(1) Applicability and Use of This Technology

(a) Liquid Injection. Liquid injection is applicable to wastes that have viscosity values sufficiently low so that the waste can be

atomized in the combustion chamber. A range of literature maximum viscosity values are reported with the low being 100 SSU and the high being 10,000 SSU. It is important to note that viscosity is temperature dependent so that while liquid injection may not be applicable to a waste at ambient conditions, it may be applicable when the waste is heated. Other factors that affect the use of liquid injection are particle size and the presence of suspended solids. Both of these waste parameters can cause plugging of the burner nozzle.

(b) Rotary Kiln/ Fluidized Bed/Fixed Hearth. These incineration technologies are applicable to a wide range of hazardous wastes. They can be used on wastes that contain high or low total organic content, high or low filterable solids, various viscosity ranges, and a range of other waste parameters. EPA has not found these technologies to be demonstrated on wastes that are comprised essentially of metals with low organic concentrations. In addition, the Agency expects that some of the high metal content wastes may not be compatible with existing and future air emission limits without emission controls far more extensive than currently practiced.

(2) Underlying Principles of Operation

(a) Liquid Injection. The basic operating principle of this incineration technology is that incoming liquid wastes are volatilized and then additional heat is supplied to the waste to destabilize the chemical bonds. Once the chemical bonds are broken, these constituents

react with oxygen to form carbon dioxide and water vapor. The energy needed to destabilize the bonds is referred to as the energy of activation.

- (b) Rotary Kiln and Fixed Hearth. There are two distinct principles of operation for these incineration technologies, one for each of the chambers involved. In the primary chamber, energy, in the form of heat, is transferred to the waste to achieve volatilization of the various organic waste constituents. During this volatilization process some of the organic constituents will oxidize to CO₂ and water vapor. In the secondary chamber, additional heat is supplied to overcome the energy requirements needed to destabilize the chemical bonds and allow the constituents to react with excess oxygen to form carbon dioxide and water vapor. The principle of operation for the secondary chamber is similar to liquid injection.
- (c) Fluidized Bed. The principle of operation for this incinerator technology is somewhat different than for rotary kiln and fixed hearth incineration, in that there is only one chamber which contains the fluidizing sand and a freeboard section above the sand. The purpose of the fluidized bed is to both volatilize the waste and combust the waste. Destruction of the waste organics can be accomplished to a better degree in this chamber than in the primary chamber of the rotary kiln and fixed hearth because of 1) improved heat transfer from fluidization of the waste using forced air and 2) the fact that the

fluidization process provides sufficient oxygen and turbulence to convert the organics to carbon dioxide and water vapor. The freeboard generally does not have an afterburner; however, additional time is provided for conversion of the organic constituents to carbon dioxide, water vapor, and hydrochloric acid if chlorine is present in the waste.

(3) Description of Incineration Technologies

- (a) Liquid Injection. The liquid injection system is capable of incinerating a wide range of gases and liquids. The combustion system has a simple design with virtually no moving parts. A burner or nozzle atomizes the liquid waste and injects it into the combustion chamber where it burns in the presence of air or oxygen. A forced draft system supplies the combustion chamber with air to provide oxygen for combustion and turbulence for mixing. The combustion chamber is usually a cylinder lined with refractory (i.e., heat resistant) brick and can be fired horizontally, vertically upward, or vertically downward. Figure 3-1 illustrates a liquid injection incineration system.
- (b) Rotary Kiln. A rotary kiln is a slowly rotating, refractory-lined cylinder that is mounted at a slight incline from the horizontal (see Figure 3-2). Solid wastes enter at the high end of the kiln, and liquid or gaseous wastes enter through atomizing nozzles in the kiln or afterburner section. Rotation of the kiln exposes the solids to the heat, vaporizes them, and allows them to combust by mixing with air. The rotation also causes the ash to move to the lower end of the kiln

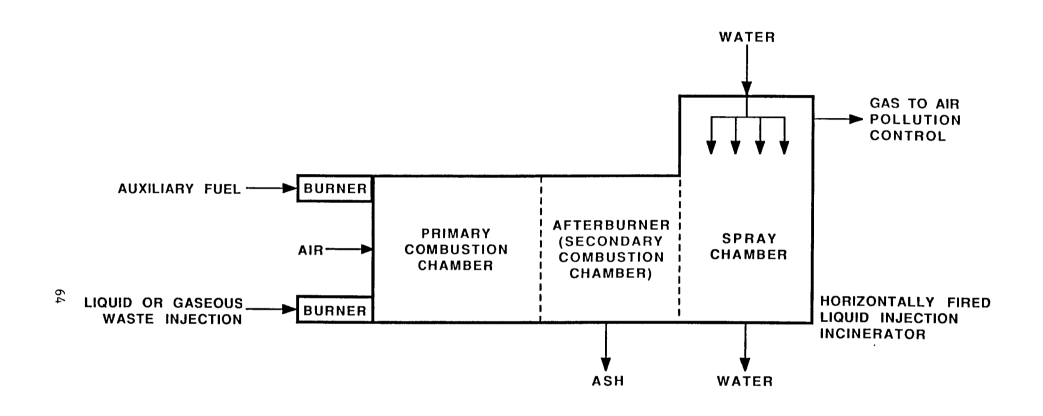


FIGURE 3-1.
LIQUID INJECTION INCINERATOR

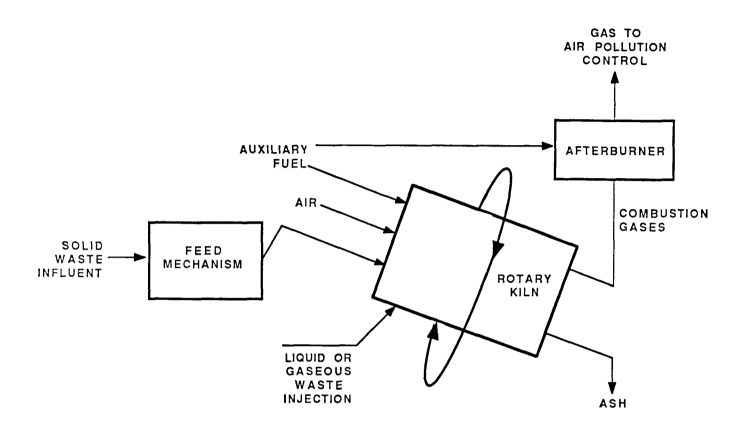


FIGURE 3-2.
ROTARY KILN INCINERATOR

where it can be removed. Rotary kiln systems usually have a secondary combustion chamber or afterburner following the kiln for further combustion of the volatilized components of solid wastes.

- (c) Fluidized Bed. A fluidized bed incinerator consists of a column containing inert particles such as sand which is referred to as the bed. Air, driven by a blower, enters the bottom of the bed to fluidize the sand. Air passage through the bed promotes rapid and uniform mixing of the injected waste material within the fluidized bed. The fluidized bed has an extremely high heat capacity (approximately three times that of flue gas at the same temperature), thereby providing a large heat reservoir. The injected waste reaches ignition temperature quickly and transfers the heat of combustion back to the bed. Continued bed agitation by the fluidizing air allows larger particles to remain suspended in the combustion zone. (See Figure 3-3)
- (d) Fixed Hearth Incineration. Fixed hearth incinerators, also called controlled air or starved air incinerators, are another major technology used for hazardous waste incineration. Fixed hearth incineration is a two-stage combustion process (see Figure 3-4). Waste is ram-fed into the first stage, or primary chamber, and burned at less than stoichiometric conditions. The resultant smoke and pyrolysis products, consisting primarily of volatile hydrocarbons and carbon monoxide, along with the normal products of combustion, pass to the secondary chamber. Here, additional air is injected to complete the

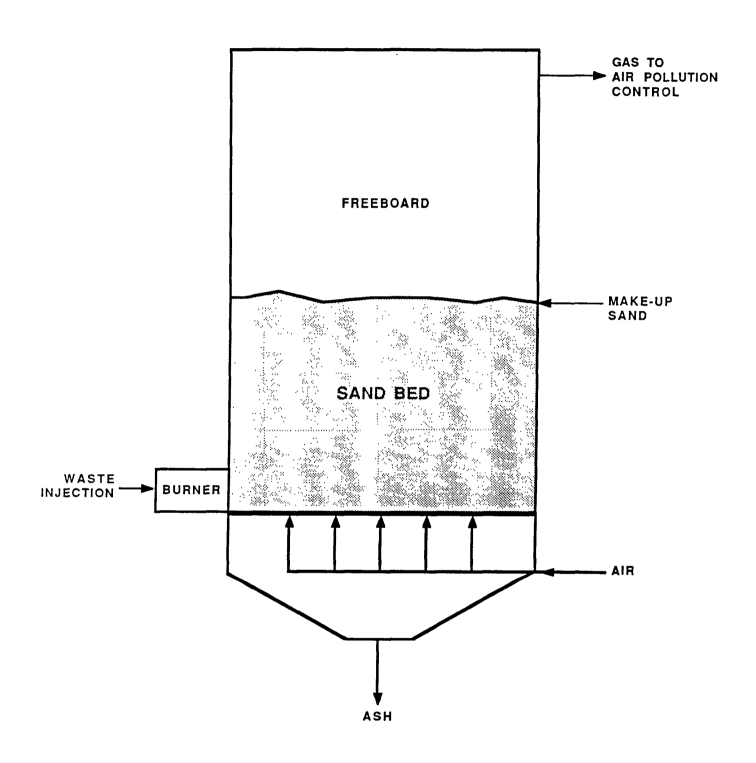


FIGURE 3-3. FLUIDIZED BED INCINERATOR

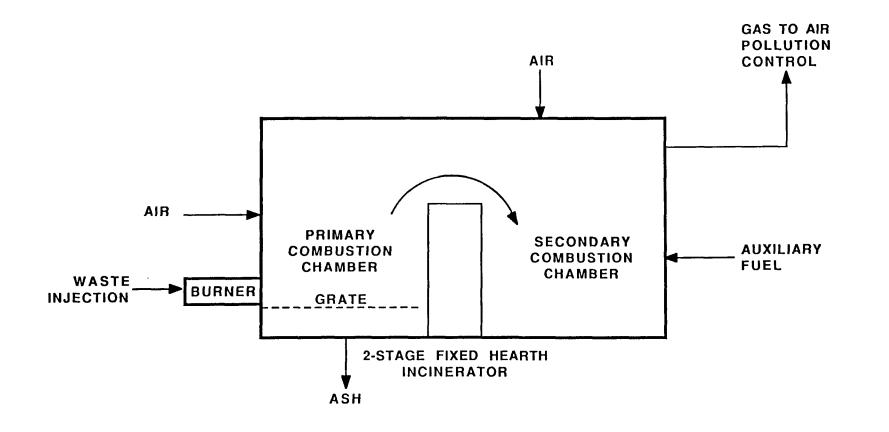


FIGURE 3-4. FIXED HEARTH INCINERATOR

combustion. This two-stage process generally yields low stack particulate and carbon monoxide (CO) emissions. The primary chamber combustion reactions and combustion gas are maintained at low levels by the starved air conditions so that particulate entrainment and carryover are minimized.

(e) Air Pollution Controls. Following incineration of hazardous wastes, combustion gases are generally further treated in an air pollution control system. The presence of chlorine or other halogens in the waste requires a scrubbing or absorption step to remover HCl and other halo-acids from the combustion gases. Ash in the waste is not destroyed in the combustion process. Depending on its composition, ash will either exit as bottom ash, at the discharge end of a kiln or hearth for example, or as particulate matter (fly ash) suspended in the combustion gas stream. Particulate emissions from most hazardous waste combustion systems generally have particle diameters less than one micron and require high efficiency collection devices to minimize air emissions. In addition, scrubber systems provide additional buffer against accidental releases of incompletely destroyed waste products due to poor combustion efficiency or combustion upsets, such as flame outs.

(4) Waste Characteristics Affecting Performance (WCAP)

(a) Liquid Injection. In determining whether liquid injection is likely to achieve the same level of performance on an untested waste as a previously tested waste, the Agency will compare dissociation bond

energies of the constituents in the untested and tested waste. This parameter is being used as a surrogate indicator of activation energy which, as discussed previously, destabilizes molecular bonds. In theory, the bond dissociation energy would be equal to the activation energy; however, in practice this is not always the case. Other energy effects (e.g., vibrational, the formation of intermediates, and interactions between different molecular bonds) may have a significant influence on activation energy.

Because of the shortcomings of bond energies in estimating activation energy, EPA analyzed other waste characteristic parameters to determine if these parameters would provide a better basis for transferring treatment standards from an untested waste to a tested waste. These parameters include heat of combustion, heat of formation, use of available kinetic data to predict activation energies, and general structural class. All of these were rejected for reasons provided below.

The heat of combustion only measures the difference in energy of the products and reactants; it does not provide information on the transition state (i.e., the energy input needed to initiate the reation). Heat of formation is used as a predictive tool for whether reactions are likely to proceed; however, there are a significant number of hazardous constituents for which these data are not available. Use of kinetic data were rejected because these data are limited and could not be used to calculate free energy values (ΔG) for the wide range of hazardous constituents to be addressed by this rule. Finally, EPA decided not to

use structural classes because the Agency believes that evaluation of bond dissociation energies allows for a more direct determination of whether a constituent will be destabilized.

- (b) Rotary Kiln/Fluidized Bed/Fixed Hearth. Unlike liquid injection, these incineration technologies also generate a residual ash. Accordingly, in determining whether these technologies are likely to achieve the same level of performance on an untested waste as a previously tested waste, EPA would need to examine the waste characteristics that affect volatilization of organics from the waste, as well as, destruction of the organics, once volatilized. Relative to volatilization, EPA will examine thermal conductivity of the entire waste and boiling point of the various constituents. As with liquid injection, EPA will examine bond energies in determining whether treatment standards for scrubber water residuals can be transferred from a tested waste to an untested waste. Below is a discussion of how EPA arrived at thermal conductivity and boiling point as the best method to assess volatilization of organics from the waste; the discussion relative to bond energies is the same for these technologies as for liquid injection and will not be repeated here.
- (i) <u>Thermal Conductivity</u>. Consistent with the underlying principles of incineration, a major factor with regard to whether a particular constituent will volatilize is the transfer of heat through the waste. In the case of rotary kiln, fluidized bed, and fixed hearth incineration,

heat is transferred through the waste by three mechanisms: radiation, convection, and conduction. For a given incinerator, heat transferred through various wastes by radiation is more a function of the design and type of incinerator than the waste being treated. Accordingly, the type of waste treated will have a minimal impact on the amount of heat transferred by radiation. With regard to convection, EPA also believes that the type of heat transfer will generally be more a function of the type and design of incinerator than the waste itself. However, EPA is examining particle size as a waste characteristic that may significantly impact the amount of heat transferred to a waste by convection and thus impact volatilization of the various organic compounds. The final type of heat transfer, conduction, is the one that EPA believes will have the greatest impact on volatilization of organic constituents. To measure this characteristic, EPA will use thermal conductivity; an explanation of this parameter, as well as, how it can be measured is provided below.

Heat flow by conduction is proportional to the temperature gradient across the material. The proportionality constant is a property of the material and referred to as the thermal conductivity. (Note: The analytical method that EPA has identified for measurement of thermal conductivity is described in Appendix D). In theory, thermal conductivity would always provide a good indication of whether a constituent in an untested waste would be treated to the same extent in

the primary incinerator chamber as the same constituent in a previously tested waste.

In practice, thermal conductivity has some limitations in assessing the transferability of treatment standards; however, EPA has not identified a parameter that can provide a better indication of heat transfer characteristics of a waste. Below is a discussion of both the limitations associated with thermal conductivity, as well as other parameters considered.

Thermal conductivity measurements, as part of a treatability comparison for two different wastes through a single incinerator, are most meaningful when applied to wastes that are homogeneous (i.e., major constituents are essentially the same). As wastes exhibit greater degrees of non-homogeneity (e.g., significant concentration of metals in soil), then thermal conductivity becomes less accurate in predicting treatability because the measurement essentially reflects heat flow through regions having the greatest conductivity (i.e., the path of least resistance) and not heat flow through all parts of the waste.

Btu value, specific heat, and ash content were also considered for predicting heat transfer characteristics. These parameters can no better account for non-homogeneity than thermal conductivity; additionally, they are not directly related to heat transfer characteristics. Therefore, these parameters do not provide a better indication of heat transfer that will occur in any specific waste.

(ii) <u>Boiling Point</u>. Once heat is transferred to a constituent within a waste, then removal of this constituent from the waste will depend on its volatility. As a surrogate of volatility, EPA is using boiling point of the constituent. Compounds with lower boiling points have higher vapor pressures and, therefore, would be more likely to vaporize. The Agency recognizes that this parameter does not take into consideration the impact of other compounds in the waste on the boiling point of a constituent in a mixture; however, the Agency is not aware of a better measure of volatility that can easily be determined.

(5) Incineration Design and Operating Parameters

(a) Liquid Injection. For a liquid injection unit, EPA's analysis of whether the unit is well designed will focus on (1) the likelihood that sufficient energy is provided to the waste to overcome the activation level for breaking molecular bonds and (2) whether sufficient oxygen is present to convert the waste constituents to carbon dioxide and water vapor. The specific design parameters that the Agency will evaluate to assess whether these conditions are met are: temperature, excess oxygen, and residence time. Below is a discussion of why EPA believes these parameters to be important, as well as a discussion of how these parameters will be monitored during operation.

It is important to point out that, relative to the development of land disposed restriction standards, EPA is only concerned with these design parameters when a quench water or scrubber water residual is generated from treatment of a particular waste. If treatment of a

particular waste in a liquid injection unit would not generate a wastewater stream, then the Agency, for purposes of land disposal treatment standards, would only be concerned with the waste characteristics that affect selection of the unit, not the above-mentioned design parameters.

(i) <u>Temperature</u>. Temperature is important in that it provides an indirect measure of the energy available (i.e., Btus/hr) to overcome the activation energy of waste constituents. As the design temperature increases, the more likely it is that the molecular bonds will be destabilized and the reaction completed.

The temperature is normally controlled automatically through the use of instrumentation which senses the temperature and automatically adjusts the amount of fuel and/or waste being fed. The temperature signal transmitted to the controller can be simultaneously transmitted to a recording device, referred to as a strip chart, and thereby continuously recorded. To fully assess the operation of the unit, it is important to know not only the exact location in the incinerator that the temperature is being monitored but also the location of the design temperature.

(ii) Excess Oxygen. It is important that the incinerator contain oxygen in excess of the stiochiometric amount necessary to convert the organic compounds to carbon dioxide and water vapor. If insufficient oxygen is present, then destabilized waste constituents could recombine to the same or other BDAT list organic compounds and potentially cause the scrubber water to contain higher concentrations of BDAT list constituents than would be the case for a well operated unit.

In practice, the amount of oxygen fed to the incinerator is controlled by continuous sampling and analysis of the stack gas. If the amount of oxygen drops below the design value, then the analyzer transmits a signal to the valve controlling the air supply and thereby increases the flow of oxygen to the afterburner. The analyzer simultaneously transmits a signal to a recording device so that the amount of excess oxygen can be continuously recorded. Again, as with temperature, it is important to know the location from which the combustion gas is being sampled.

- (iii) <u>Carbon Monoxide</u>. Carbon monoxide is an important operating parameter because it provides an indication of the extent to which the waste organic constituents are being converted to CO₂ and water vapor. As the carbon monoxide level increases, it indicates that greater amounts of organic waste constituents are unreacted or partially reacted. Increased carbon monoxide levels can result from insufficient excess oxygen, insufficient turbulence in the combustion zone, or insufficient residence time.
- (iv) <u>Waste Feed Rate</u>. The waste feed rate is important to monitor because it is correlated to the residence time. The residence time is associated with a specific Btu energy value of the feed and a specific volume of combustion gas generated. Prior to incineration, the Btu value of the waste is determined through the use of a laboratory device known as a bomb calorimeter. The volume of combustion gas generated from the waste to be incinerated is determined from an analysis referred to as an

ultimate analysis. This analysis determines the amount of elemental constituents present which include carbon, hydrogen, sulfur, oxygen, nitrogen, and halogens. Using this analysis plus the total amount of air added, the volume of combustion gas can be calculated. Having determined both the Btu content and the expected combustion gas volume, the feed rate can be fixed at the desired residence time. Continuous monitoring of the feed rate will determine whether the unit was operated at a rate corresponding to the designed residence time.

(b)Rotary Kiln. For this incineration, EPA will examine both the primary and secondary chamber in evaluating the design of a particular incinerator. Relative to the primary chamber, EPA's assessment of design will focus on whether it is likely that sufficient energy will be provided to the waste in order to volatilize the waste constituents. For the secondary chamber,

analogous to the sole liquid injection incineration chamber, EPA will examine the same parameters discussed previously under liquid injection incineration. These parameters will not be discussed again here.

The particular design parameters to be evaluated for the primary chamber are: kiln temperature, residence time, and revolutions per minute. Below is a discussion of why EPA believes these parameters to be important, as well as a discussion of how these parameters will be monitored during operation.

(i) <u>Temperature</u>. The primary chamber temperature is important, in that it provides an indirect measure of the energy input (i.e., BTUs/hr)

that is available for heating the waste. The higher the temperature is designed to be in a given kiln, the more likely it is that the constituents will volatilize. As discussed earlier under "Liquid Injection", temperature should be continuously monitored and recorded. Additionally, it is important to know the location of the temperature sensing device in the kiln.

- (ii) Residence Time. This parameter is important in that it affects whether sufficient heat is transferred to a particular constituent in order for volatilization to occur. As the time that the waste is in the kiln is increased, a greater quantity of heat is transferred to the hazardous waste constituents. The residence time will be a function of the specific configuration of the rotary kiln including the length and diameter of the kiln, the waste feed rate, and the rate of rotation.
- (iii) Revolutions Per Minute (RPM). This parameter provides an indication of the turbulence that occurs in the primary chamber of a rotary kiln. As the turbulence increases, the quantity of heat transferred to the waste would also be expected to increase. However, as the RPM value increases, the residence time decreases resulting in a reduction of the quantity of heat transferred to the waste. This parameter needs to be carefully evaluated because it provides a balance between turbulence and residence time.
- (c) Fluidized Bed. As discussed previously, in the section on "Underlying Principles of Operation", the primary chamber accounts for almost all of the conversion of organic wastes to carbon dioxide, water

vapor, and acid gas if halogens are present. The secondary chamber will generally provide additional residence time for thermal oxidation of the waste constituents. Relative to the primary chamber, the parameters that the Agency will examine in assessing the effectiveness of the design are temperature, residence time, and bed pressure differential. The first two were discussed under rotary kiln and will not be discussed here. The latter, bed pressure differential, is important in that it provides an indication of the amount of turbulence and, therefore, indirectly the amount of heat supplied to the waste. In general, as the pressure drop increases, both the turbulence and heat supplied increase. The pressure drop through the bed should be continuously monitored and recorded to ensure that the designed valued is achieved.

(d) Fixed Hearth. The design considerations for this incineration unit are similar to a rotary kiln with the exception that rate of rotation (i.e., RPMs) is not an applicable design parameter. For the primary chamber of this unit, the parameters that the Agency will examine in assessing how well the unit is designed are the same as discussed under rotary kiln; for the secondary chamber (i.e., afterburner), the design and operating parameters of concern are the same as previously discussed under "Liquid Injection".

3.2.2 Fuel Substitution

Fuel substitution involves using hazardous waste as a fuel in industrial furnaces or in boilers for generation of steam. The hazardous waste may be blended with other nonhazardous wastes (e.g., municipal sludge) and/or fossil fuels.

(1) Applicability and Use of This Technology

Fuel substitution has been used with industrial waste solvents, refinery wastes, synthetic fibers/petrochemical wastes and waste oils. It can also be used when combusting other waste types produced during the manufacturing of pharmaceuticals, pulp and paper and pesticides. These wastes can be handled in a solid, liquid or gaseous form.

The most common types of units in which waste fuels are burned are industrial furnaces and industrial boilers. Industrial furnaces include a diverse variety of industrial processes that produce heat and/or products by burning fuels. They include blast furnaces, smelters, and coke ovens. Industrial boilers are units wherein fuel is used to produce steam for process and plant use. Industrial boilers typically use coal, oil, or gas as the primary fuel source.

There are a number of parameters that affect the selection of fuel substitution. These are:

- Halogen content of the waste.
- Inorganic solids content (ash content) of the waste, particularly heavy metals.
- · Heating value of the waste.
- · Viscosity of the waste (for liquids).
- Filterable solids concentration (for liquids).
- Sulfur content.

If halogenated organics are burned, halogenated acids and free halogen are among the products of combustion. These released corrosive

gases may require subsequent treatment prior to venting to the atmosphere. Also, halogens and halogenated acids formed during combustion are likely to severely corrode boiler tubes and other process equipment. For this reason, halogenated wastes are blended into fuels only at very low concentrations to minimize such problems. High chlorine content can also lead to the incidental production (at very low concentrations) of other hazardous compounds such as PCBs (polychlorinated biphenyls), PCDDs (chlorinated dibenzo-p-dioxins), PCDFs (chlorinated dibenzofurans), and chlorinated phenols.

High inorganic solids content (i.e., ash content) of wastes may cause two problems: (1) scaling in the boiler, and (2) particulate air emissions. Scaling results from deposition of inorganic solids on the walls of the boiler. Particulate emissions are produced by noncombustible inorganic constituents that flow out of the boiler with the gaseous combustion products. Due to these problems, wastes with significant concentrations of inorganic materials are not usually handled in boilers unless they have an air pollution control system.

Industrial furnaces vary in their tolerance to inorganic constituents. Heavy metal concentrations, found in both halogenated and nonhalogenated wastes used as fuel, can cause environmental concern because they may be emitted either in the gaseous emissions from the combustion process, in the ash residues, or in any produced solids. The partitioning of the heavy metals to these residual streams primarily depends on the volatility of the metal, waste matrix, and furnace design.

The heating value of the waste must be sufficiently high (either alone or in combination with other fuels) to maintain combustion temperatures consistent with efficient waste destruction and operation of the boiler or furnace. For many applications, only supplemental fuels having minimum heating values of 4,400 to 5,600 kcal/kg (8,000 to 10,000 BTU/lb) are considered to be feasible. Below this value, the unblended fuel would not be likely to maintain a stable flame and its combustion would release insufficient energy to provide needed steam generation potential in the boiler, or the necessary heat for an industrial furnace. Some wastes with heating values of less than 4,400 kcal/kg (8,000 BTU/lb) can be used if sufficient auxiliary fuel is employed to support combustion or if special designs are incorporated into the combustion device. Occasionally, for wastes with heating values higher than virgin fuels, blending with auxiliary fuel may be required to prevent overheating or over charging the combustion device.

In combustion devices designed to burn liquid fuels, the viscosity of liquid waste must be low enough so that it can be atomized in the combustion chamber. If viscosity is too high, heating of storage tanks may be required prior to combustion. For atomization of liquids, a viscosity of 165 centistokes (750 Saybolt Seconds Universal (SSU)) or less is typically required.

If filterable material suspended in the liquid fuel prevents or hinders pumping or atomization, it will be unacceptable.

Sulfur content in the waste may prevent burning of the waste due to potential atmospheric emissions of sulfur oxides. For instance there are proposed Federal sulfur oxide emission regulations for certain new source industrial boilers (51 FR 22385). Air pollution control devices are available to remove sulfur oxides from the stack gases.

(2) <u>Underlying Principles of Operation</u>

For a boiler and most industrial furnaces there are two distinct principles of operation. Initially, energy in the form of heat is transferred to the waste to achieve volatilization of the various waste constituents. For liquids, volatilization energy may also be supplied by using pressurized atomization. The energy used to pressurize the liquid waste allows the atomized waste to break into smaller particles thus enhancing its rate of volatilization. The volatilized constituents then require additional energy to destabilize the chemical bonds and allow the constituents to react with oxygen to form carbon dioxide and water vapor. The energy needed to destabilize the chemical bonds is referred to as the energy of activation.

(3) Physical Description of the Process

As stated, there are a number of industrial applications that can use fuel substitution. Therefore, there is no one process description that will fit all of these applications. However, the following section provides a general description of industrial kilns (one form of industrial furnace) and industrial boilers.

(a) Kilns. Combustible wastes have the potential to be used as fuel in kilns and, for waste liquids, are often used with oil to co-fire kilns. Coal-fired kilns are capable of handling some solid wastes. In the case of cement kilns, there are usually no residuals requiring land disposal since any ash formed becomes part of the product or is removed by particulate collection systems and recycled back to the kiln. The only residuals may be low levels of unburned gases escaping with combustion products. If this is the case, air pollution control devices may be required.

Three types of kilns are particularly applicable: cement kilns, lime kilns, and light-weight aggregate kilns.

(i) <u>Cement kilns</u>. The cement kiln is a rotary furnace which is a refractory-lined steel shell used to calcine a mixture of calcium, silicon, aluminum, iron, and magnesium-containing minerals. The kiln is normally fired by coal or oil. Liquid and solid combustible wastes may then serve as auxiliary fuel. Temperatures within the kiln are typically between 1,380°C and 1,540°C (2,500°F to 2,800°F). To date, only liquid hazardous wastes have been burned in cement kilns.

Most cement kilns have a dry particulate collection device (i.e., either an electrostatic precipitator or baghouse) with the collected fly ash recycled back to the kiln. Build up of metals or other noncombustibles is prevented through their incorporation in the product cement. Many types of cement require a source of chloride so that most halogenated liquid hazardous wastes currently can be burned in cement kilns. Available information shows that scrubbers are not used.

(ii) <u>Lime kilns</u>. Quick-lime (CaO) is manufactured in a calcination process using limestone (CaCO $_3$) or dolomite (CaCO $_3$ and MgCO $_3$). These raw materials are also heated in a refractory-lined rotary kiln, typically to temperatures of 980°C to 1,260°C (1,800°F to 2,300°F). Lime kilns are less likely to burn hazardous wastes than cement kilns because product lime is often added to potable water systems. Only one lime kiln currently burns hazardous waste in the U.S. That particular facility sells its product lime for use as flux or as refractory in blast furnaces.

As with cement kilns, any collected fly ash is recycled back to the lime kiln, resulting in no residual streams from the kiln. Available information shows that scrubbers are not used.

(iii) <u>Lightweight aggregate kilns</u>. Lightweight aggregate kilns heat clay to produce an expanded lightweight inorganic material used in portland cement formulations and other applications. The kiln has a normal temperature range of 1,100°C to 1,150°C (2,000°F to 2,100°F). Lightweight aggregate kilns are less amenable to combustion of hazardous wastes as fuels than other kilns described above due to their lack of material in the kiln to adsorb halogens. As a result burning of halogenated organics in these kilns would likely require afterburners to ensure complete destruction of the halogenated organics and scrubbers to control acid gas production. Such controls would produce a wastewater residual stream subject to treatment standards.

(b) Industrial Boilers. A boiler is a closed vessel in which water is transformed into steam by the application of heat. Normally, heat is supplied by the combustion of pulverized coal, fuel oil, or gas. These fuels are fired into a combustion chamber with nozzles and burners that provide mixing with air. Liquid wastes, and granulated solid wastes in the case of grate-fired boilers, can be burned as auxiliary fuel in a boiler. Few grate-fired boilers burn hazardous wastes, however. For liquid-fired boilers, residuals requiring land disposal are only generated when the boiler is shut down and cleaned. This is generally done once or twice per year. Other residuals from liquid-fired boilers would be the gas emission stream which would consist of any products of incomplete combustion, along with the normal combustion products. For example, chlorinated wastes would produce acid gases. If this is the case, air pollution control devices may be required. For solid fired boilers, an ash normally is generated. This ash may contain residual amounts of organics from the blended waste/fuels as well as noncombustible materials. Land disposal of this ash would require compliance with applicable BDAT treatment standards.

(4) Waste Characteristics Affecting Performance

For cement kilns and lime kilns, and lightweight aggregate kilns burning nonhalogenated wastes (i.e., no scrubber is needed to control acid gases), no residual waste streams would be produced. Any noncombustible material in the waste would leave the kiln in the product stream. As a result, in transferring standards EPA would not examine

waste characteristics affecting performance but rather would determine the applicability of fuel substitution. That is, EPA would investigate the parameters affecting treatment selection. For kilns these parameters (as mentioned previously) are Btu content, percent filterable solids, halogenated organics content, viscosity, and sulfur content.

Lightweight aggregate kilns burning halogenated organics and boilers burning wastes containing any noncombustibles will produce residual streams subject to treatment standards. In determining whether fuel substitution is likely to achieve the same level of performance on an untreated waste as a previously treated waste, EPA will examine:

(a) relative volatility of the waste constituents, (b) the heat transfer characteristics (for solids); and (c) the activation energy for combustion.

(a) Relative volatility. The term relative volatility (α) refers to the ease with which a substance present in a solid or liquid waste will vaporize from that waste upon application of heat from an external source. Hence, it bears a relationship to the equilibrium vapor pressure of the substance.

EPA recognizes that the relative volatilities can not be measured or calculated directly for the types of wastes generally treated in an industrial boiler or furnace. The Agency believes that the best measure of relative volatility is the boiling point of the various hazardous constituents and will, therefore, use this parameter in assessing volatility of the organic constituents.

(b) Heat transfer characteristics. Consistent with the underlying principles of combustion in aggregate kilns or boilers, a major factor with regard to whether a particular constituent will volatilize is the transfer of heat through the waste. In the case of industrial boilers burning solid fuels, heat is transferred through the waste by three mechanisms: radiation, convection, and conduction. For a given boiler it can be assumed that the type of waste will have a minimal impact on the heat transferred from radiation. With regard to convection, EPA believes that the range of wastes treated would exhibit similar properties with regard to the amount of heat transferred by convection. Therefore, EPA will not evaluate radiation convection heat transfer properties of wastes in determining similar treatability. For solids, the third heat transfer mechanism, conductivity, is the one principally operative or most likely to change between wastes. The method that EPA believes can be used to determine thermal conductivity of a nonwastewater can be found in Appendix D.

Using thermal conductivity measurements as part of a treatability comparison for two different wastes through a given boiler or furnace is most meaningful when applied to wastes that are homogeneous. As wastes exhibit greater degrees of non-homogeneity, then thermal conductivity becomes less accurate in predicting treatability because the measurement essentially reflects heat flow through regions having the greatest conductivity (i.e., the path of least resistance and not heat flow through all parts of the waste). Nevertheless, EPA has not identified a

better alternative to thermal conductivity, even for wastes that are non-homogeneous.

Other parameters considered for predicting heat transfer characteristics were Btu value, specific heat, and ash content. These parameters can neither better account for non-homogeneity nor better predict heat transferability through the waste.

(c) Activation energy. Given an excess of oxygen, an organic waste in an industrial furnace or boiler would be expected to convert to CO₂ and H₂O provided that the activation energy is achieved. Activation energy is the quantity of heat (energy) needed to destabilize molecular bonds and create reactive intermediates so that the oxidation (combustion) reaction will proceed to completion. As a measure of activation energy, EPA is using bond dissociation energies. In theory, the bond dissociation energy would be equal to the activation energy; however, in practice this is not always the case.

In some instances, bond energies will not be available and will have to be estimated or other energy effects (e.g., vibrational) and other reactions will have a significant influence on activation energy. Because of the shortcomings of bond energies in estimating activation energy, EPA analyzed other waste characteristic parameters to determine if these parameters would provide a better basis for transferring treatment standards from an untested waste to a tested waste. These parameters included heat of combustion, heat of formation, use of available kinetic data to predict activation energies, and general structural class. All of these were rejected for reasons provided below.

The heat of combustion only measures the difference in energy of the products and reactants; it does not provide information on the transition state (i.e., the energy input needed to initiate the reaction). Heat of formation is used as a predictive tool for whether reactions are likely to proceed; however, there are a significant number of hazardous constituents for which these data are not available. Use of available kinetic data were rejected because while it could be used to calculate some free energy values (ΔG), it could not be used for the wide range of hazardous constituents. Finally, EPA decided not to use structural classes because the Agency believes that evaluation of bond dissociation energies allows for a more direct comparison.

(5) <u>Design and Operating Parameters</u>

(a) Design Parameters. Cement kilns and lime kilns, along with aggregate kilns burning nonhalogenated wastes produce no residual streams. Their design and operation is such that any wastes that are incompletely destroyed will be contained in the product. As a result, the Agency will not look at design and operating values for such devices since treatment, per se, cannot be measured through detection of constituents in residual streams. In this instance it is important merely to ensure the waste is appropriate for combustion in the kilns and the kiln is operated in a manner that will produce a useable product.

Specifically, cement, lime, and aggregate kilns are only demonstrated on liquid hazardous wastes. Such wastes must be sufficiently free of filterable solids to avoid plugging the burners at the hot end of the

kiln. Viscosity also must be low enough to inject the waste into the kiln through the burners. The sulfur content is not a concern unless the concentration in the waste is sufficiently high as to exceed federal, state, or local air pollution standards promulgated for industrial boilers.

The design parameters which normally affect the operation of an industrial boiler (and aggregate kilns with residual streams) with respect to hazardous waste treatment are (i) the design temperature, (ii) the design retention time of the waste in the combustion chamber, and (iii) turbulence in the combustion chamber. Evaluation of these parameters would be important in determining if an industrial boiler or industrial furnace is adequately designed for effective treatment of hazardous wastes. The rationale for selection of three parameters is given below.

(i) <u>Design temperature</u>. Industrial boilers are generally designed based on their steam generation potential (BTU output). This factor is related to the design combustion temperature, which in turn depends on the amount of fuel burned, and its BTU value. The fuel feed rates and combustion temperatures of industrial boilers are generally fixed based on the BTU values of fuels normally handled (e.g., No. 2 versus No. 6 fuel oils). When wastes are to be blended with fossil fuels for combustion, the blending, based on BTU values, must be such that the resulting BTU value of the mixture is close to that of the fuel value used in design of the boiler. Industrial furnaces also are designed to

operate at specific ranges of temperature in order to produce the desired product (e.g., lightweight aggregate). The blended waste/fuel mixture should be capable of maintaining the design temperature range.

- (ii) <u>Retention time</u>. A sufficient retention time of combustion products is normally necessary to ensure that the hazardous substances being combusted (or formed during combustion) are completely oxidized. Retention times on the order of a few seconds are normally needed at normal operating conditions. For industrial furnaces, as well as boilers, the retention time is a function of the size of the furnace and the fuel feed rates. For most boilers and furnaces the retention time usually exceeds a few seconds.
- (iii) <u>Turbulence</u>. Boilers are designed so that fuel and air are intimately mixed. This helps ensure that complete combustion takes place. The shape of the boiler, and the method of fuel and air feed influence the turbulence required for good mixing. Industrial furnaces also are designed for turbulent mixing where fuel and air are mixed.
- (b) Operating Parameters. The operating parameters which normally affect the performance of an industrial boiler and many industrial furnaces with respect to treatment of hazardous wastes are (i) air flow rate, (ii) fuel feed rate, (iii) steam pressure or rate of production, and (iv) temperature. EPA believes that these four parameters will be used to determine if an industrial boiler burning blended fuels containing hazardous waste constituents is properly operated. The rationale for selection of these four operating parameters is given

- below. Most industrial furnaces will monitor similar parameters, but some exceptions are noted below.
- (i) Air feed rate. An important operating parameter in boilers and many industrial furnaces is the oxygen content in the flue gas which is a function of the air feed rate. Stable combustion of a fuel generally occurs within a specific range of air-to-fuel ratios. An oxygen analyzer in the combustion gases can be used to control the feed ratio of air to fuel to assure complete thermal destruction of the waste and efficient operation of the boiler. When necessary, the air flow rate can be increased or decreased to maintain proper fuel to oxygen ratios. Some industrial furnaces do not completely combust fuels (e.g., coke ovens and blast furnaces) hence oxygen concentration in the flue gas is a meaningless variable.
- (ii) <u>Fuel feed rate</u>. The rate at which fuel is injected into the boiler or industrial furnace will determine the thermal output of the system per unit of time (BTU/hr). If steam is produced, steam pressure monitoring will indirectly determine if the fuel feed rate is adequate. However, various velocity and mass measurement devices can be used to monitor fuel flow directly.
- (iii) <u>Steam pressure or rate of production</u>. Steam pressure in boilers provides a direct measure of the thermal output of the system and is directly monitored by use of in-system pressure gauges. Increases or decreases in steam pressure can be effected by increasing or decreasing the fuel and air feed rates within certain operating design limits. Most

industrial furnaces do not produce steam, but instead a product (e.g., cement, aggregate) and monitor the rate of production.

(iv) <u>Temperature</u>. Temperatures are monitored and controlled in industrial boilers to assure the quality and flow rate of steam.

Therefore, complex monitoring systems are frequently installed in the combustion unit to provide a direct reading of temperature. The efficiency of combustion in industrial boilers is dependent on combustion temperatures. Temperature may be adjusted to design settings by increasing or decreasing air and fuel feed rate.

Wastes should not be added to primary fuels until the boiler temperature reaches the minimum needed for destruction of the wastes.

Temperature instrumentation and control should be designed to stop waste addition in the event of process upsets.

Monitoring and control of temperature in industrial furnaces are also critical to the product quality; e.g., lime, cement, or aggregate kilns, that require minimum operating temperatures. Kilns have very high thermal inertia in the refractory and in-process product, high residence times, and high air flow rates, so that even in the case of a momentary stoppage of fuel flow to the kiln, organic constituents are likely to continue to be destroyed. The main operational control required for wastes burned in kilns is to stop waste flow in the event of low kiln temperature, loss of the electrical power to the combustion air fan, and loss of primary fuel flow.

(c) Other Operating Parameters. In addition to the four operating parameters discussed above, EPA considered and then discarded one additional parameter. Fuel to waste blending ratios were also considered. However, while the blending is done to yield a uniform BTU content fuel, blending ratios will vary on a wide range dependent on the BTU content of the wastes and fuels being used.

3.2.3 Stabilization of Metals

Stabilization refers to a broad class of treatment processes that chemically reduce the mobility of hazardous constituents in a waste.

Solidification and fixation are other terms that are sometimes used synonymously for stabilization or to describe specific variations within the broader class of stabilization. Related technologies are encapsulation and thermoplastic binding; however, EPA considers these technologies to be distinct from stabilization in that the operational principles are significantly different.

(1) Applicability and Use of This Technology

Stabilization is used when a waste contains metals that will leach from the waste when it is contacted by water. In general, this technology is applicable to wastes containing BDAT list metals, having a high filterable solids content, low TOC content, and low oil and grease content. This technology is commonly used to treat residuals generated from treatment of electroplating wastewaters. For some wastes, an alternative to stabilization is metal recovery.

(2) Underlying Principles of Operation

The basic principle underlying this technology is that stabilizing agents and other chemicals are added to a waste in order to minimize the amount of metal that leaches. The reduced leachability is accomplished by the formation of a lattice structure and/or chemical bonds that bind the metals to the solid matrix and, thereby, limit the amount of metal constituents that can be leached when water or a mild acid solution comes into contact with the waste material.

There are two principal stabilization processes used; these are cement based and lime based. A brief discussion of each is provided below. In both cement-based or lime/pozzolan-based techniques, the stabilizing process can be modified through the use of additives, such as silicates, that control curing rates or enhance the properties of the solid material.

(a) Portland Cement-Based Process. Portland cement is a mixture of powdered oxides of calcium, silica, aluminum, and iron, produced by kiln burning of materials rich in calcium and silica at high temperatures (i.e., 1400°C to 1500°C). When the anhydrous cement powder is mixed with water, hydration occurs and the cement begins to set. The chemistry involved is complex because many different reactions occur depending on the composition of the cement mixture.

As the cement begins to set, a colloidal gel of indefinite composition and structure is formed. Over a period of time, the gel swells and forms a matrix composed of interlacing, thin, densely-packed

silicate fibrils. Constituents present in the waste slurry (e.g., hydroxides and carbonates of various heavy metals, are incorporated into the interstices of the cement matrix. The high pH of the cement mixture tends to keep metals in the form of insoluble hydroxide and carbonate salts.) It has been hypothesized that metal ions may also be incorporated into the crystal structure of the cement matrix, but this hypothesis has not been verified.

(b) Lime/Pozzolan-Based Process. Pozzolan, which contains finely divided, noncrystalline silica (e.g., fly ash or components of cement kiln dust), is a material that is not cementitious in itself, but becomes so upon the addition of lime. Metals in the waste are converted to silicates or hydroxides which inhibit leaching. Additives, again, can be used to reduce permeability and thereby further decrease leaching potential.

(3) <u>Description of Stabilization Processes</u>

In most stabilization processes, the waste, stabilizing agent, and other additives, if used, are mixed and then pumped to a curing vessel or area and allowed to cure. The actual operation (equipment requirements and process sequencing) will depend on several factors such as the nature of the waste, the quantity of the waste, the location of the waste in relation to the disposal site, the particular stabilization formulation to be used, and the curing rate. After curing, the solid formed is recovered from the processing equipment and shipped for final disposal.

In instances where waste contained in a lagoon is to be treated, the material should be first transferred to mixing vessels where stabilizing agents are added. The mixed material is then fed to a curing pad or vessel. After curing, the solid formed is removed for disposal. Equipment commonly used also includes facilities to store waste and chemical additives. Pumps can be used to transfer liquid or light sludge wastes to the mixing pits and pumpable uncured wastes to the curing site. Stabilized wastes are then removed to a final disposal site.

Commercial concrete mixing and handling equipment generally can be used with wastes. Weighing conveyors, metering cement hoppers, and mixers similar to concrete batching plants have been adapted in some operations. Where extremely dangerous materials are being treated, remote-control and in-drum mixing equipment, such as that used with nuclear waste, can be employed.

(4) Waste Characteristics Affecting Performance

In determining whether stabilization is likely to achieve the same level of performance on an untested waste as on a previously tested waste, the Agency will focus on the characteristics that inhibit the formation of either the chemical bonds or the lattice structure. The four characteristics EPA has identified as affecting treatment performance are the presence of (a) fine particulates, (b) oil and grease, (c) organic compounds, and (d) certain inorganic compounds.

(a) Fine Particulates. For both cement-based and lime/pozzolan-based processes, the literature states that very fine solid

materials (i.e., those that pass through a No. 200 mesh sieve, 74 um particle size) can weaken the bonding between waste particles and cement by coating the particles. This coating can inhibit chemical bond formation and decreases the resistance of the material to leaching.

- (b) Oil and Grease. The presence of oil and grease in both cement-based and lime/pozzolan-based systems results in the coating of waste particles and the weakening of the bonding between the particle and the stabilizing agent. This coating can inhibit chemical bond formation and thereby, decrease the resistance of the material to leaching.
- (c) Organic Compounds. The presence of organic compounds in the waste interferes with the chemical reactions and bond formation which inhibit curing of the stabilized material. This results in a stabilized waste having decreased resistance to leaching.
- (d) Sulfate and Chlorides. The presence of certain inorganic compounds will interfere with the chemical reactions, weakening bond strength and prolonging setting and curing time. Sulfate and chloride compounds may reduce the dimensional stability of the cured matrix, thereby increasing leachability potential.

Accordingly, EPA will examine these constituents when making decisions regarding transfer of treatment standards based on stabilization.

(5) <u>Design and Operating Parameters</u>

In designing a stabilization system, the principal parameters that are important to optimize so that the amount of leachable metal

constituents is minimized are (a) selection of stabilizing agents and other additives, (b) ratio of waste to stabilizing agents and other additives, (c) degree of mixing, and (d) curing conditions.

(a) Selection of stabilizing agents and other additives. The stabilizing agent and additives used will determine the chemistry and structure of the stabilized material and, therefore, will affect the leachability of the solid material. Stabilizing agents and additives must be carefully selected based on the chemical and physical characteristics of the waste to be stabilized. For example, the amount of sulfates in a waste must be considered when a choice is being made between a lime/pozzolan and a Portland cement-based system.

In order to select the type of stabilizing agents and additives, the waste should be tested in the laboratory with a variety of materials to determine the best combination.

(b) Amount of stabilizing agents and additives. The amount of stabilizing agents and additives is a critical parameter in that sufficient stabilizing materials are necessary in the mixture to bind the waste constituents of concern properly, thereby making them less susceptible to leaching. The appropriate weight ratios of waste to stabilizing agent and other additives are established empirically by setting up a series of laboratory tests that allow separate leachate - testing of different mix ratios. The ratio of water to stabilizing agent (including water in waste) will also impact the strength and leaching characteristics of the stabilized material. Too much water will cause

low strength; too little will make mixing difficult and, more importantly, may not allow the chemical reactions that bind the hazardous constituents to be fully completed.

- (c) Mixing. The conditions of mixing include the type and duration of mixing. Mixing is necessary to ensure homogeneous distribution of the waste and the stabilizing agents. Both undermixing and overmixing are undesirable. The first condition results in a nonhomogeneous mixture; therefore, areas will exist within the waste where waste particles are neither chemically bonded to the stabilizing agent nor physically held within the lattice structure. Overmixing, on the other hand, may inhibit gel formation and ion adsorption in some stabilization systems. As with the relative amounts of waste, stabilizing agent, and additives within the system, optimal mixing conditions generally are determined through laboratory tests. During treatment it is important to monitor the degree (i.e., type and duration) of mixing to ensure that it reflects design conditions.
- (d) Curing conditions. The curing conditions include the duration of curing and the ambient curing conditions (temperature and humidity). The duration of curing is a critical parameter to ensure that the waste particles have had sufficient time in which to form stable chemical bonds and/or lattice structures. The time necessary for complete stabilization depends upon the waste type and the stabilization used. The performance of the stabilized waste (i.e., the levels of constituents in the leachate) will be highly dependent upon whether complete stabilization

has occurred. Higher temperatures and lower humidity increase the rate of curing by increasing the rate of evaporation of water from the solidification mixtures. However, if temperatures are too high, the evaporation rate can be excessive and result in too little water being available for completion of the stabilization reaction. The duration of the curing process should also be determined during the design stage and typically will be between 7 and 28 days.

3.2.4 Chemical Precipitation

(1) Applicability and Use of This Technology

Chemical precipitation is used when dissolved metals are to be removed from solution. This technology can be applied to a wide range of wastewaters containing dissolved BDAT list metals and other metals as well. This treatment process has been practiced widely by industrial facilities since the 1940s.

(2) Underlying Principles of Operation

The underlying principle of chemical precipitation is that metals in wastewater are removed by the addition of a treatment chemical that converts the dissolved metal to a metal precipitate. This precipitate is less soluble than the original metal compound, and therefore settles out of solution, leaving a lower concentration of the metal present in the solution. The principal chemicals used to convert soluble metal compounds to the less soluble forms include: lime $(Ca(OH)_2)$, caustic (NaOH), sodium sulfide (Na_2S) , and, to a lesser extent, soda ash (Na_2CO_3) , phosphate, and ferrous sulfide (FeS).

The solubility of a particular compound will depend on the extent to which the electrostatic forces holding the ions of the compound together can be overcome. The solubility will change significantly with temperature; most metal compounds are more soluble as the temperature increases. Additionally, the solubility will be affected by the other constituents present in a waste. As a general rule, nitrates, chlorides, and sulfates are more soluble than hydroxides, sulfides, carbonates, and phosphates.

An important concept related to treatment of the soluble metal compounds is pH. This term provides a measure of the extent to which a solution contains either an excess of hydrogen or hydroxide ions. The pH scale ranges from 0 to 14; with 0 being the most acidic, 14 representing the highest alkalinity or hydroxide ion (OH⁻) content, and 7.0 being neutral.

When hydroxide is used, as is often the case, to precipitate the soluble metal compounds, the pH is frequently monitored to ensure that sufficient treatment chemicals are added. It is important to point out that pH is not a good measure of treatment chemical addition for compounds other than hydroxides; when sulfide is used, for example, facilities might use an oxidation-reduction potential meter (ORP) correlation to ensure that sufficient treatment chemical is used.

Following conversion of the relatively soluble metal compounds to metal precipitates, the effectiveness of chemical precipitation is a function of the physical removal, which usually relies on a settling

process. A particle of a specific size, shape, and composition will settle at a specific velocity, as described by Stokes' Law. For a batch system, Stokes' law is a good predictor of settling time because the pertinent particle parameters remain essentially constant. Nevertheless, in practice, settling time for a batch system is normally determined by empirical testing. For a continuous system, the theory of settling is complicated by factors such as turbulence, short-circuting, and velocity gradients, increasing the importance of the empirical tests.

(3) <u>Description of the Technology</u>

The equipment and instrumentation required for chemical precipitation varies depending on whether the system is batch or continuous. Both operations are discussed below; a schematic of the continuous system is shown in Figure 3-5.

For a batch system, chemical precipitation requires only a feed system for the treatment chemicals and a second tank where the waste can be treated and allowed to settle. When lime is used, it is usually added to the reaction tank in a slurry form. In a batch system, the supernate is usually analyzed before discharge, thus minimizing the need for instrumentation.

In a continuous system, additional tanks are necessary, as well as instrumentation to ensure that the system is operating properly. In this system, the first tank that the wastewater enters is referred to as an equalization tank. This is where the waste can be mixed in order to provide more uniformity, minimizing wide swings in the type and

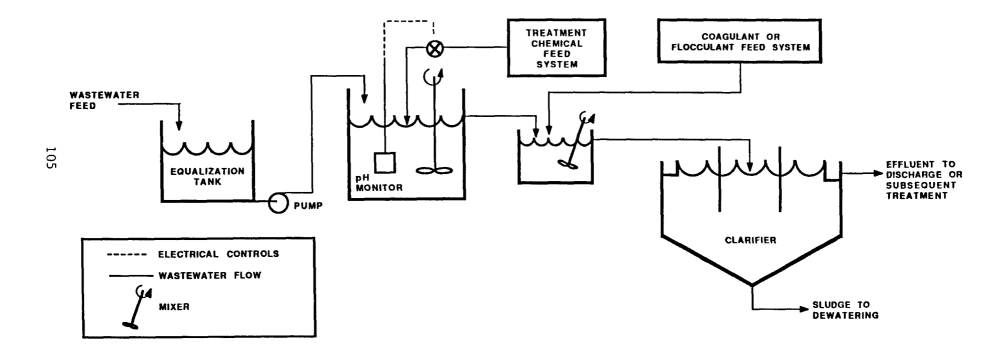


FIGURE 3-5. CONTINUOUS CHEMICAL PRECIPITATION

concentration of constituents being sent to the reaction tank. It is important to reduce the variability of the waste sent to the reaction tank because control systems inherently are limited with regard to the maximum fluctuations that can be managed.

Following equalization, the waste is pumped to a reaction tank where treatment chemicals are added; this is done automatically by using instrumentation that senses the pH of the system and then pneumatically adjusts the position of the treatment chemical feed valve such that the design pH value is achieved. Both the complexity and the effectiveness of the automatic control system will vary depending on the variation in the waste and the pH range that is needed to properly treat the waste.

An important aspect of the reaction tank design is that it be well-mixed so that the waste and the treatment chemicals are both dispersed throughout the tank, in order to ensure comingling of the reactant and the treatment chemicals. In addition, effective dispersion of the treatment chemicals throughout the tank is necessary to properly monitor and, thereby, control the amount of treatment chemicals added.

After the waste is reacted with the treatment chemical, it flows to a quiescent tank where the precipitate is allowed to settle and subsequently be removed. Settling can be chemically assisted through the use of flocculating compounds. Flocculants increase the particicle size and density of the precipitated solids, both of which increase the rate of settling. The particular flocculating agent that will best improve settling characteristics will vary depending on the particular waste;

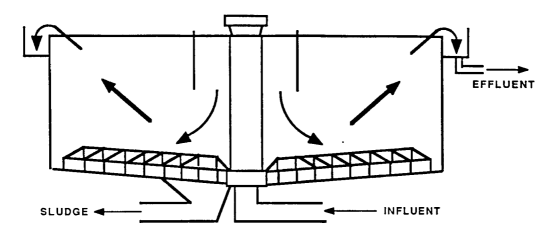
selection of the flocculating agent is generally accomplished by performing laboratory bench tests. Settling can be conducted in a large tank by relying solely on gravity or be mechanically assisted through the use of a circular clarifier or an inclinded separator. Schematics of the latter two separators are shown in Figures 3-6 and 3-7.

Filtration can be used for further removal of precipitated residuals both in cases where the settling system is underdesigned and in cases where the particles are difficult to settle. Polishing filtration is discussed in a separate technology section.

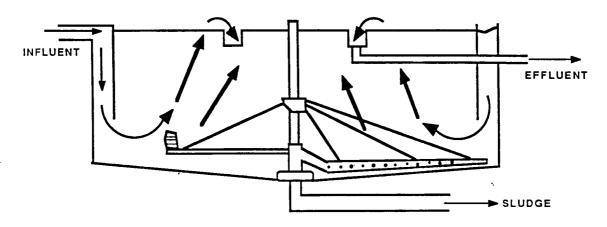
(4) Waste Characteristics Affecting Performance

In determining whether chemical preciptation is likely to achieve the same level of performance on an untested waste as a previously tested waste, we will examine the following waste characteristics: (a) the concentration and type of the metal(s) in the waste, (b) the concentration of suspended solids (TSS), (c) the concentration of dissolved solids (TDS), (d) whether the metal exists in the wastewater as a complex, and (e) the oil and grease content. These parameters either affect the chemical reaction of the metal compound, the solubility of the metal precipitate, or the ability of the precipitated compound to settle.

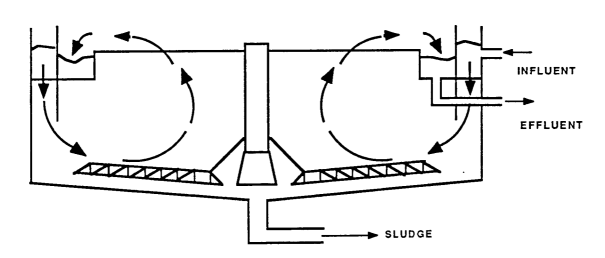
(a) Concentration and type of metals. For most metals, there is a specific pH at which the metal hydroxide is least soluble. As a result, when a waste contains a mixture of many metals, it is not possible to operate a treatment system at a single pH which is optimal for the removal of all metals. The extent to which this affects treatment depends on the particular metals to be removed, and their



CENTER FEED CLARIFIER WITH SCRAPER SLUDGE REMOVAL SUSTEM



RIM FEED - CENTER TAKEOFF CLARIFIER WITH HYDRAULIC SUCTION SLUDGE REMOVAL SYSTEM



RIM FEED - RIM TAKEOFF CLARIFIER

FIGURE 3-6 CIRCULAR CLARIFIERS 108

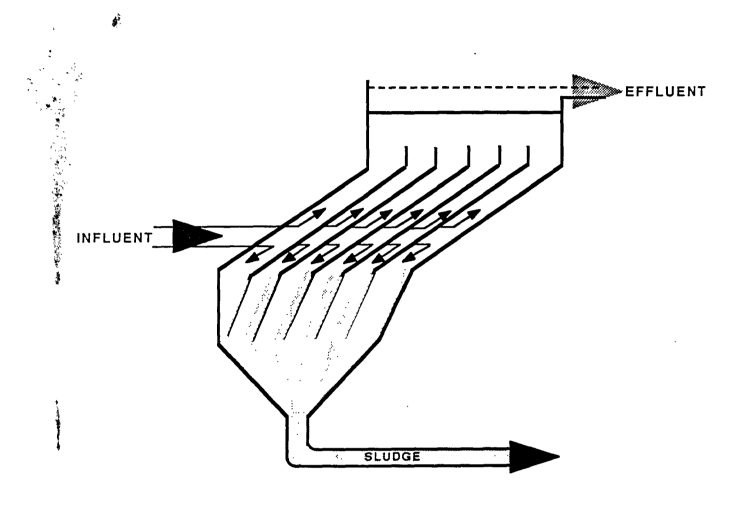


FIGURE 3-7
INCLINED PLATE SETTLER

concentrations. An alternative can be to operate multiple precipitations, with intermediate settling, when the optimum pH occurs at markedly different levels for the metals present. The individual metals and their concentrations can be measured using EPA Method 6010.

- (b) Concentration and type of total suspended solids (TSS).

 Certain suspended solid compounds are difficult to settle because of either their particle size or shape. Accordingly, EPA will evaluate this characteristic in assessing transfer of treatment performance. Total suspended solids can be measured by EPA Wastewater Test Method 160.2.
- (c) Concentration of total dissolved solids (TDS). Available information shows that total dissolved solids can inhibit settling. The literature states that poor flocculation is a consequence of high TDS and shows that higher concentrations of total suspended solids are found in treated residuals. Poor flocculation can adversely affect the degree to which precipitated particles are removed. Total dissolved solids can be measured by EPA Wastewater Test Method 160.1.
- (d) Complexed metals. Metal complexes consist of a metal ion surrounded by a group of other inorganic or organic ions or molecules (often called ligands). In the complexed form, the metals have a greater solubility and, therefore, may not be as effectively removed from solution by chemical precipitation. EPA does not have an analytical method to determine the amount of complexed metals in the waste. The Agency believes that the best measure of complexed metals is to analyze for some common complexing compounds (or complexing agents) generally

found in wastewater for which analytical methods are available. These complexing agents include ammonia, cyanide, and EDTA. The analytical method for cyanide is EPA Method 9010. The method for EDTA is ASTM Method D3113. Ammonia can be analyzed using EPA Wastewater Test Method 350.

(e) Oil and grease content. The oil and grease content of a particular waste directly inhibits the settling of the precipitate. Suspended oil droplets float in water and tend to suspend particles such as chemical precipitates that would otherwise settle out of the solution. Even with the use of coagulants or flocculants, the separation of the precipitate is less effective. Oil and grease content can be measured by EPA Method 9071.

(5) Design and Operating Parameters

The parameters that EPA will evaluate when determining whether a chemical precipitation system is well designed are: (a) design value for treated metal concentrations, as well as other characteristics of the waste used for design purposes (e.g., total suspended solids), (b) pH, (c) residence time, (d) choice of treatment chemical, (e) choice of coagulant/flocculant, and (f) mixing. Below is an explanation of why EPA believes these parameters are important to a design analysis; in addition, EPA explains why other design criteria are not included in EPA's analysis.

(a) Treated and untreated design concentrations. EPA pays close attention to the treated concentration the system is designed to achieve

when determining whether to sample a particular facility. Since the system will seldom out-perform its design, EPA must evaluate whether the design is consistent with best demonstrated practice.

The untreated concentrations that the system is designed to treat are important in evaluating any treatment system. Operation of a chemical precipitation treatment system with untreated waste concentrations in excess of design values can easily result in poor performance.

- (b) pH. The pH is important, because it can indicate that sufficient treatment chemical (e.g., lime) is added to convert the metal constituents in the untreated waste to forms that will precipitate. The pH also affects the solubility of metal hydroxides and sulfides, and therefore directly impacts the effectiveness of removal. In practice, the design pH is determined by empirical bench testing, often referred to as "jar" testing. The temperature at which the "jar" testing is conducted is important in that it also affects the solubility of the metal precipitates. Operation of a treatment system at temperatures above the design temperature can result in poor performance. In assessing the operation of a chemical precipitation system, EPA prefers continuous data on the pH and periodic temperature conditions throughout the treatment period.
- (c) Residence time. The residence time is important because it impacts the completeness of the chemical reaction to form the metal precipitate and, to a greater extent, amount of precipitate that settles out of solution. In practice, it is determined by "jar" testing. For

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- (c) Residence time. The residence time is important because it impacts the completeness of the chemical reaction to form the metal precipitate and, to a greater extent, amount of precipitate that settles out of solution. In practice, it is determined by "jar" testing. For

continuous systems, EPA will monitor the feed rate to ensure that the system is operated at design conditions. For batch systems, EPA will want information on the design parameter used to determine sufficient settling time (e.g., total suspended solids).

- (d) Choice of treatment chemical. A choice must be made as to what type of precipitating agent (i.e., treatment chemical) will be used. The factor that most affects this choice is the type of metal constituents to be treated. Other design parameters, such as pH, residence time, and choice of coagulant/flocculant agents, are based on the selection of the treatment chemical.
- (e) Choice of coagulant/flocculant. This is important because these compounds improve the settling rate of the precipitated metals and allows for smaller systems (i.e., lower retention time) to achieve the same degree of settling as a much larger system. In practice, the choice of the best agent and the required amount is determined by "jar" testing.
- (f) Mixing. The degree of mixing is a complex assessment which includes, among other things, the energy supplied, the time the material is mixed, and the related turbulence effects of the specific size and shape of the tank. EPA will, however, consider whether mixing is provided and whether the type of mixing device is one that could be expected to achieve uniform mixing. For example, EPA may not use data from a chemical precipitation treatment system where an air hose was placed in a large tank to achieve mixing.

3.2.5 Sludge Filtration

(1) Applicability and Use of This Technology

Sludge filtration, also known as sludge dewatering or cake-formation filtration, is a technology used on wastes that contain high concentrations of suspended solids, generally higher than one percent. The remainder of the waste is essentially water. Sludge filtration is applied to sludges, typically those that have settled to the bottom of clarifiers, for dewatering. After filtration, these sludges can be dewatered to 20 to 50 percent solids.

(2) <u>Underlying Principle of Operation</u>

The basic principle of filtration is the separation of particles from a mixture of fluids and particles by a medium that permits the flow of the fluid but retains the particles. As would be expected, larger particles are easier to separate from the fluid than smaller particles. Extremely small particles, in the colloidal range, may not be filtered effectively and may appear in the treated waste. To mitigate this problem, the wastewater should be treated prior to filtration to modify the particle size distribution in favor of the larger particles, by the use of appropriate precipitants, coagulants, flocculants, and filter aids. The selection of the appropriate precipitant or coagulant is important because it affects the particles formed. For example, lime neutralization usually produces larger, less gelatinous particles than does caustic soda precipitation. For larger particles that become too small to filter effectively because of poor resistance to shearing, shear

resistance can be improved by the use of coagulants and flocculants.

Also, if pumps are used to feed the filter, shear can be minimized by designing for a lower pump speed, or by use of a low shear type of pump.

(3) Technology Description

For sludge filtration, settled sludge is either pumped through a cloth-type filter media (such as in a plate and frame filter that allows solid "cake" to build up on the media) or the sludge is drawn by vacuum through the cloth media (such as on a drum or vacuum filter, which also allows the solids to build). In both cases the solids themselves act as a filter for subsequent solids removal. For a plate and frame type filter, removal of the solids is accomplished by taking the unit off line, opening the filter and scraping the solids off. For the vacuum type filter, cake is removed continuously. For a specific sludge, the plate and frame type filter will usually produce a drier cake than a vacuum filter. Other types of sludge filters, such as belt filters, are also used for effective sludge dewatering.

(4) Waste Characteristics Affecting Performance

The following characteristics of the waste will affect performance of a sludge filtration unit:

- size of particles and
- type of particles.
- (a) Size of particles. The smaller the particle size, the more the particles tend to go through the filter media. This is especially true for a vacuum filter. For a pressure filter (like a plate and frame),

smaller particles may require higher pressures for equivalent throughput, since the smaller pore spaces between particles create resistance to flow.

(b) Type of particles. Some solids formed during metal precipitation are gelatinous in nature and cannot be dewatered well by cake-formation filtration. In fact, for vacuum filtration a cake may not form at all. In most cases solids can be made less gelatinous by use of the appropriate coagulants and coagulant dosage prior to clarification, or after clarification but prior to filtration. In addition, the use of lime instead of caustic soda in metal precipitation will reduce the formation of gelatinous solids. Also the addition of filter aids to a gelatinous sludge, such as lime or diatomaceous earth, will help significantly. Finally, precoating the filter with diatomaceous earth prior to sludge filtration will assist in dewatering gelatinous sludges.

(5) Design and Operating Parameters

For sludge filtration, the following design and operating variables affect performance:

- type of filter selected,
- size of filter selected,
- feed pressure, and
- use of coagulants or filter aids.
- (a) Type of filter. Typically, pressure type filters (such as a plate and frame) will yield a drier cake than a vacuum type filter and will also be more tolerant of variations in influent sludge characteristics. Pressure type filters, however, are batch operations,

so that when cake is built up to the maximum depth physically possible (constrained by filter geometry), or to the maximum design pressure, the filter is turned off while the cake is removed. A vacuum filter is a continuous device (i.e., cake discharges continuously), but will usually be much larger than a pressure filter with the same capacity. A hybrid device is a belt filter, which mechanically squeezes sludge between two continuous fabric belts.

- (b) Size of filter. As with in-depth filters, the larger the filter, the greater its hydraulic capacity and the longer the filter runs between cake discharge.
- (c) Feed pressure. This parameter impacts both the design pore size of the filter and the design flow rate. It is important that in treating waste that the design feed pressure not be exceeded, otherwise particles may be forced through the filter medium resulting in ineffective treatment.
- (c) Use of coagulants. Coagulants and filter aids may be mixed with filter feed prior to filtration. Their effect is particularly significant for vacuum filtration in that it may make the difference in a vacuum filter between no cake and a relatively dry cake. In a pressure filter, coagulants and filter aids will also significantly improve hydraulic capacity and cake dryness. Filter aids, such as diatomaceous earth, can be precoated on filters (vacuum or pressure) for particularly difficult to filter sludges. The precoat layer acts somewhat like an in-depth filter in that sludge solids are trapped in the precoat pore

spaces. Use of precoats and most coagulants or filter aids significantly increases the amount of sludge solids to be disposed of. However, polyelectrolyte coagulant usage usually does not increase sludge volume significantly because the dosage is low.

3.3 Performance Data

3.3.1 BDAT List Organics Treatment Data

The Agency collected nine data sets (untreated and treated waste data) to characterize the treatment performance of rotary kiln incineration on KOO1. Six of these data sets are from KOO1 wastes from wood preservation processes using creosote based preservative chemicals and three are from KOO1 wastes containing pentachlorophenol. The data presented include BDAT list volatile, semivolatile, and metal constituents detected in the untreated KOO1, the ash (nonwastewater residual), and scrubber water (wastewater residual) from rotary kiln incineration. Tables 3-1 through 3-9 present the nine data sets for the BDAT list constituents detected in the untreated and treated waste samples from rotary kiln incineration. Operating data collected during the incineration test burns are presented in Tables 3-10 to 3-13.

3.3.2 BDAT List Metals Treatment Data

The Agency does not have performance data specifically for treatment of the BDAT list metals in the ash and scrubber water generated from rotary kiln incineration of KOO1. However, EPA does have treatment performance data on wastes that the Agency believes are sufficiently

similar to these residuals with regard to parameters affecting treatment selection. In these cases, the treatment performance data for BDAT list metals in similar wastes were available for transfer for the development of BDAT treatment standards for KOO1.

- (1) <u>Wastewater residuals</u>. The performance data that the Agency has for wastewaters include 11 data sets from the Onsite Engineering Report for Envirite Corporation. We believe these data can be used to transfer levels of performance because they contain the constituents of concern in concentrations at least as high as the concentrations expected to be in K001 scrubber water. These data are presented in Table 3-14.
- (2) Nonwastewater residuals. The performance data for stabilization of K001 nonwastewater residuals were transferred from stabilization of F006. EPA examined all available treatment performance data from wastes that are considered to be similar with regard to the parameters affecting treatment selection. Based on the BDAT list metals present in the untreated waste and the treated residuals, as well as the waste characteristics of the residuals treatment data were identified from wastes similar to K001 treated scrubber water residuals and incinerator ash. F006 are the wastewater treatment sludges from electroplating which the Agency believes to be similar to the nonwastewaters from treatment of K001 wastewaters and incinerator ash. These treatment performance data are presented in Tables 3-15 to 3-17. The BDAT list metals present in K001 nonwastewater residuals are generally lower in concentration than

the BDAT list metals in F006. Further, the metals in K001 residuals are likely to be present in the oxide form since they have resulted from an incineration process. Metals in F006 are typically in the hydroxide form. The Agency believes that metals in the form of oxides are more readily immobilized (less leachable) than metals in the form of hydroxide. For these reasons the Agency believes that treatment standards for F006 can be attained even more readily for K001 nonwastewater residuals.

Table 3-1 Rotary Kiln Incineration of K001 - Creosote $\\ \text{Sample Set No} \quad 1$

BDAT List Constituent	Untreatedwaste(ppb)	Treated nonwastewater(ash) Total(ppb)	Treated wastewater (scrubber water) (ug/1)
Volatile Organics			
Benzene Toluene Ethyl benzene	56 110 57	<50 <50 <50	<50 <50 <50
Xylenes	120	<50	<50
			
	Untreatedwaste(ppm)	Treated nonwastewater(ash) Total (ppm)	Treated wastewater <u>(scrubber water)</u> (ug/l)
Semivolatile Organi	waste (ppm)	nonwastewater (ash) Total	wastewater (scrubber water)
Acenaphthalene	waste(ppm)	nonwastewater (ash) Total (ppm)	wastewater (scrubber water) (ug/1)
Acenaphtha lene Acenaphthene	. waste (ppm) cs <4600 21,000	nonwastewater(ash)	wastewater (scrubber water) (ug/l) <20 <10
Acenaphthalene Acenaphthene Anthracene	<pre>. waste</pre>	nonwastewater(ash)	wastewater (scrubber water) (ug/l) <20 <10 <10
Acenaphthalene Acenaphthene Anthracene Chrysene	<pre>. waste</pre>	nonwastewater(ash)	<pre>wastewater (scrubber water) (ug/l) <20 <10 <10 <15</pre>
Acenaphthalene Acenaphthene Anthracene Chrysene Fluorene	<pre></pre>	nonwastewater (ash) Tota; (ppm) <1 15 <0 65 <0 65 <0 85 <0 65	<pre>wastewater (scrubber water) (ug/l) <20 <10 <10 <15 <10</pre>
Acenaphthalene Acenaphthene Anthracene Chrysene Fluorene Naphthalene	<pre></pre>	nonwastewater (ash) Total (ppm) <1 15 <0 65 <0 65 <0 85 <0 65 <0 65 <0 55	<pre>wastewater (scrubber water) (ug/l) <20 <10 <10 <15 <10 <10 <10<<10<<10<<10<<10<<10<<10<<10</pre>
Acenaphthalene Acenaphthene Anthracene Chrysene Fluorene	<pre></pre>	nonwastewater (ash) Tota; (ppm) <1 15 <0 65 <0 65 <0 85 <0 65	<pre>wastewater (scrubber water) (ug/l) <20 <10 <10 <15 <10</pre>

Table 3-1 (continued)

Sample Set No 1

	Untreated	Treat	Treated	
BDAT List	waste	<u>nonwastewater (ash)</u> Total TCLP		wastewater (scrubber water
Constituent	(ppm)	(ppm)	(mg/l)	(mg/1)
Metals				
Antimony	<17	<17	0 035	< 0 170
Arsenic	2 6	5.3	< 0 020	0 041
Barıum	63	81	0 38	0 76
Beryllium	< 0 5	<0.5	<0.005	0 002
Cadmium	3.4	<2.0	0.020	0 87
Chromium	5 0	6 1	< 0 035	0 35
Copper	35	86	0.020	0 42
Mercury	0.35	<1 25	< 0 002	5 0 016
Nickel	2 1	3 6	< 0 075	0 48
Lead	170	<21	< 0 21	2 5
Selenium	1 5	2 3	<0.020	0.057
Silver	<3 5	< 3 5	< 0 035	< 0 035
Thallium	7 7	<2 5	< 0 025	< 0 028
Vanadıum	<4 0	4 4	< 0 040	0.040
Zinc	170	1 9	0 020	5 0

Reference Onsite Engineering Report for K001-Creosote

Table 3-2 Rotary Kiln Incineration of KOO1 - Creosote

Sample Set No. 2

BDAT List Constituent	Untreated waste (ppb)	Treated nonwastewater(ash)	Treated wastewater (scrubber water) (ug/l)
Volatile Organics			
Benzene	€0	< 50	<50
Toluene	120	< 50	< 50
Ethyl benzene	5€	< 50	< 50
Xylenes	130	<50	<50
	Untreated <u>waste</u> (ppm)	Treated nonwastewater (ash) Total (ppm)	Treated wastewater (scrubber water) (ug/l)
Semivolatile Organi	waste (ppm)	nonwastewater(as") Total	wastewater (scrubber water)
<u>Semivolatile Organi</u> Acenaphthalene	waste (ppm)	nonwastewater(as") Total	wastewater (scrubber water)
	waste (ppm)	nonwastewater (ash) Total (ppm)	wastewater <u>(scrubber water)</u> (ug/l)
Acenaphthalene	waste (ppm)	nonwastewater (as**) Total (ppm) <1.15 <0.65 <0.65	wastewater (scrubber water) (ug/l)
Acenaphthalene Acenaphthene	waste(ppm)	nonwastewater (asn) Total (ppm) <1.15 <0.65 <0.65 <0.85	<pre>vastewater (scrubber water) (ug/1) <20 <10 <10 <15</pre>
Acenaphthalene Acenaphthene Anthracene	waste (ppm)	nonwastewater (as**) Total (ppm) <1.15 <0.65 <0.65 <0.85 <0.65	<pre>vastewater (scrubber water) (ug/l) <20 <10 <10 <15 <10</pre>
Acenaphthalene Acenaphthene Anthracene Chrysene	waste (ppm)	nonwastewater (as**) Total (ppm) <1.15 <0.65 <0.65 <0.85 <0.65 <0.55	<pre>wastewater (scrubber_water) (ug/1) <20 <10 <10 <15</pre>
Acenaphthalene Acenaphthene Anthracene Chrysene Fluorene	1000 15,000 7300 4200 12,000 40,000 32,000	nonwastewater (as**) Total (ppm) <1.15 <0.65 <0.65 <0.85 <0.65 <0.55 <1.80	<pre>vastewater (scrubber water) (ug/l) <20 <10 <10 <15 <10</pre>
Acenaphthalene Acenaphthene Anthracene Chrysene Fluorene Naphthalene	1000 15,000 7300 4200 12,000 40,000	nonwastewater (as**) Total (ppm) <1.15 <0.65 <0.65 <0.85 <0.65 <0.55	<pre>vastewater (scrubber water) (ug/l) <20 <10 <10 <15 <10 <10 <10<</pre>

Table 3-2 (continued)
Sample Set No 2

		Trea	Treated wastewater		
DDAT	Untreated		nonwastewater (ash)		
BDAT List	<u>waste</u>	Total	TCLP	(scrubber water)	
Constituent	(ppm)	(ppm)	(mg/l)	(mg/l)	
<u>Metals</u>					
Ant imony	<17	<17	< 0 17	< 0 170	
Arsenic	<2 0	5 3	< 0 02	0 15	
Barıum	58	74	0 57	0.80	
Beryllium	< 0 5	< 0 5	< 0.005	<0.005	
Cadmium	5 4	<2.0	< 0 02	<0.99	
Chromium	4 8	5 3	< 0 035	0 74	
Copper	32	100	0 030	0 50	
Mercury	0 35	<1 25	< 0 002	5 0 060	
Nickel	\7 5	4. ნ	< 0 075	0.51	
Lead	160	<21	<0.21	4 5	
Selenium	1 4	1.9	< 0 02	0 11	
Silver	<3 5	< 3 5	< 0 035	0 010	
Thallium	8 0	<2.5	< 0 025	2.4	
Vanadium	<4 0	4 1	< 0 04	0.060	
Zinc	170	1 8	0 050	7.1	

Reference. Onsite Engineering Report for K001-Creosote

BDAT List Constituent	Untreated <u>waste</u> (ppb)	Treated nonwastewater (ash) Total (ppb)	Treated wastewater <u>(scrubber water)</u> (ug/l)
Volatile Organics			
Benzene	61	<50	<50
Toluene	100	< 50	< 50
Ethyl benzene	55	< 50	< 50
Xylenes	120	< 50	<50
	Untreated	Treated nonwastewater (ash)	Treated wastewater
	waste	Total	(scrubber water)
	(ppm)	(ppm)	(-ug/1)
	ıcs		,
<u>Semivolatile Organ</u>			
Semivolatile Organ Acenaphthalene	<4600	<1 15	<20
		<1 15 <0.65	<20 <10
Acenaphthalene	<4600		
Acenaphthalene Acenaphthene	<4600 19,000	<0.65	<10
Acenaphthalene Acenaphthene Anthracene	<4600 19,000 12,000	<0.65 <0.65	<10 <10
Acenaphthalene Acenaphthene Anthracene Chrysene	<4600 19,000 12,000 4800	<0.65 <0.65 <0.85 <0.65 <0.55	<10 <10 <15 <10 <10
Acenaphthalene Acenaphthene Anthracene Chrysene Fluorene	<4600 19,000 12,000 4800 16,000	<0.65 <0.65 <0.85 <0.65 <0.55 <1.80	<10 <10 <15 <10
Acenaphthalene Acenaphthene Anthracene Chrysene Fluorene Naphthalene	<4600 19,000 12,000 4800 16,000	<0.65 <0.65 <0.85 <0.65 <0.55	<10 <10 <15 <10 <10

Table 3-3 (continued)
Sample Set No 3

	Untreated	Trea	ated vater (ash)	Treated wastewater
BDAT List	waste	Total	TCLP	(scrubber water)
Constituent	(ppm)	(ppm)	(mg/1)	(mg/1)
<u>Metals</u>				
Ant imony	<17	<17	0 040	< 0 17
Arsenic	2 1	9.0	<0.02	0 16
Barıum	70	61	0.53	0 56
Beryllıum	< 0 5	<0.5	<0.005	0.001
Cadmium	3 1	< 2 0	< 0 02	0 95
Chromium	6 4	5.8	< 0 035	0 60
Copper	39	98	0.030	0 43
Mercury	<1 2	5 <1 25	<0.002	5 0.008
Nickel	2 1	5 0	0 020	0 56
Lead	150	<21	< 0 21	3 5
Selenium	1 2	1.8	<0.020	0 090
Silver	<3 5	<3.5	<0.035	0 010
Thallium	6 8	<2 5	< 0 025	2 8
Vanadium	<4 0	4.1	< 0 040	0 040
Zinc	160	3.2	0.080	9 1

Reference. Onsite Engineering Report for K001-Creosote.

Table 3-4 Rotary Kilm Incineration of K001 - Creosote

Sample Set No 4

BDAT List Constituent	Untreated waste (ppb)	Treated nonwastewater (ash) Total (ppb)	Treated wastewater (scrubber water) (ug/l)
Volatile Organics			
Benzene	51	< 50	<50
Toluene	110	< 50	< 50
Ethyl benzene	72	< 50	< 50
Xylenes	130	< 50	<50
	Untreated <u>waste</u> (ppm)	Treated nonwastewater(ash) Total(ppm)	Treated wastewater <u>(scrubper water)</u> (ug/l)
Semivolatile Organ	<u>108</u>		
Acenaphtha lene	<4600	<1 15	<20
Acenaphthene	16,000	< 0 65	<10
Anthracene	8500	< 0 65	<10
Anthracene			
	4100	<0 85	<15
	4100 14.000	< 0 65	<10
Chrysene	4100 14.000 32.000	<0 65 <0 55	<10 <10
Chrysene Fluorene Naphthalene Phenanthrene	4100 14,000 32,000 29,000	<0 65 <0 55 <1 80	<10 <10 <30
Chrysene Fluorene Naphthalene	4100 14.000 32.000	<0 65 <0 55	<10 <10

Table 3-4 (continued)
Sample Set No 4

	Untreated	Treat nonwastewa	Treated wastewater		
BDAT List	waste	Total	TCLP	(scrubber w	ater)
Constituent	(ppm)	(mqq)	(mg/l)	(mg/1)	
<u>Metals</u>					
Antimony	<17	<17	0.040	0	040
Arsenic	2 5	10	< 0 020	0	28
Barıum	59	48	0 31	1	0
Beryllium	< 0 5	< 0 5	< 0 005	0	001
Cadmium	2 4	< 2 0	< 0 020	1	2
Chromium	7 0	8 7	< 0 035	1	0
Copper	39	110	< 0 030	0	51
Mercury	0 40	<1 25	< 0 0025	0	29
Nickel	2.8	5.2	<0.075	0	60
Lead	110	<21	~0.210	5	4
Selenium	1 1	2 5	< 0 020	0	12
Silver	<3.5	< 3 5	< 0 035	0	020
Thallium	5 3	<2 5	< 0 025	ā	4
Vanadıum	0 62	4.2	< 0 040	0	080
Zinc	120	2 8	0 010	11	

Reference Onsite Engineering Report for KOC1-Creosote.

Table 3-5 Rotary Kiln Incineration of K001 - Creosote $Sample \ \mbox{Set No.} \ 5$

BDAT List Constituent	Untreatedwaste(ppb)	Treated nonwastewater(ash) Total (ppb)	Treated wastewater (scrubber water) (ug/})	
Volatile Organics				
Benzene	58	<50	< 50	
Toluene	110	< 50	<50	
Ethyl benzene	71	<50	<50	
Xylenes	130	<50	<50	
	Untreated waste (ppm)	Treated nonwastewater (ash) Total (ppm)	Treated wastewater (scrubber water) (ug/l)	
Semivolatile Organia	<u> </u>			
Semivolatile Organia Acemaphthalene	<u>cs</u> <4600	<1 15	<20	
-		< 0 65	<10	
Acenaphthalene	<4600	< 0.65 < 0.65	<10 <10	
Acemaphthalene Acenaphthene Anthracene Chrysene	<4600 19,000 7400 4200	<0.65 <0.65 <0.85	<10 <10 <15	
Acenaphthalene Acenaphthene Anthracene Chrysene Fluorene	<4600 19,000 7400 4200 16,000	<0.65 <0.65 <0.85 <0.65	<10 <10 <15 <10	
Acenaphthalene Acenaphthene Anthracene Chrysene Fluorene Naphthalene	<4600 19,000 7400 4200 16,000 29,000	<0.65 <0.65 <0.85 <0.65 <0.55	<10 <10 <15 <10 <10	
Acenaphthene Anthracene Chrysene	<4600 19,000 7400 4200 16,000	<0.65 <0.65 <0.85 <0.65	<10 <10 <15 <10	

Table 3-5 (continued)
Sample Set No 5

	Untreated	Treat		Treated
BDAT List		nonwastewa Total	TCLP	wastewater
Constituent	<u>waste</u> (ppm)	(ppm)	(mg/1)	(scrubber water) (mg/l)
	(ρρπ)	(pp)	(mg/ · /	(11197-17
<u>Metals</u>				
Ant imorry	<17	<17	0 040	< 0 170
Arsenic	0 7	13	0.025	0 25
Barium	12	56	0.22	0 90
Beryllium	< 0 5	<0.5	< 0 005	0.002
Cadmium	0.79	<2 0	< 0 020	0 45
Chromium	16	10	< 0 035	0 65
Copper	12	130	0 008	0.45
Mercury	0 79	<1.25	< 0 002	5 0 19
Nickel	1 8	€ 8	< 0 075	0 70
Lead	37	<21	<0.210	3.3
Selenium	0 3	26	0 004	0 038
Silver	<3 5	<3.5	< 0 035	0 010
Thallium	2.2	<25	< 0 025	3.8
Variadium	<4.0	5 <i>2</i>	0 020	0.050
Zinc	40	3 0	0 030	8.2

Reference: Onsite Engineering Report for KOOl-Creosote

Table 3-6 Rotary Kilm Incineration of K001 - Creosote

Sample Set No 6

BDAT List Constituent	Untreated <u>waste</u> (ppb)	Treated nonwastewater (ash) Total (ppb)	Treated wastewater (scrubber water) (ug/l)
Volatile Organics			
Benzene	83	<50	< 50
To luene	170	< 50	< 50
Ethyl benzene	87	< 50	< 50
Xyl e nes	170	< 50	<50
		Treated	
	Untreated wiste (ppm)	nonwastewater (ash) Total (ppm)	Treated wastewater (scrubber water) (ug/l)
Semivolatile Organi	wiste (ppm)	(ash) Total	wastewater (scrubber water)
Semivolatile Organi Acenaphthalene	wiste (ppm)	(ash) Total	wastewater (scrubber water)
Acenapnthalene Acenapnthene		(ash) Total (ppm)	wastewater (scrubber water) (ug/l)
Acenaphthalene		(ash) Total (ppm) <1 15 <0 65 <0 65	wastewater (scrubber water) (ug/l) <20 <10 <10
Acenapnthalene Acenapnthene Anthracene Chrysene	- 4600 17,000 9100 4300	(ash) Total (ppm) <1 15 <0 65 <0 65 <0 85	<pre>wastewater (scrubber_water) (ug/l) <20 <10 <10 <15</pre>
Acenaphthalene Acenaphthene Anthracene Chrysene Fluorene	- 4600 17,000 9100 4300 14,000	(ash) Total (ppm) <1 15 <0 65 <0 65 <0 85 <0 65	<pre>wastewater (scrubber water) (ug/l) <20 <10 <10 <15 <10</pre>
Acenapnthalene Acenapnthene Anthracene Chrysene Fluorene Naphthalene	- 4600 17,000 9100 4300 14,000 45,000	(ash) Total (ppm) <1 15 <0 05 <0 65 <0 85 <0 65 <0 65 <0 55	<pre>wastewater (scrubber water) (ug/l) <20 <10 <10 <15 <10 <10 <10<<10</pre>
Acenaphthalene Acenaphthene Anthracene Chrysene Fluorene Naphthalene Phenanthrene	4600 17,000 9100 4300 14,000 45,000 36,000	(ash) Total (ppm) <1 15 <0 65 <0 65 <0 65 <0 65 <1 80	<pre>wastewater {scrubber water} (ug/l) <20 <10 <10 <15 <10 <10 <30</pre>
Acenapnthalene Acenapnthene Anthracene Chrysene Fluorene Naphthalene	- 4600 17,000 9100 4300 14,000 45,000	(ash) Total (ppm) <1 15 <0 05 <0 65 <0 85 <0 65 <0 65 <0 55	<pre>wastewater (scrubber water) (ug/l) <20 <10 <10 <15 <10 <10 <10<<10</pre>

Table 3-6 (continued)

Sample Set No 6

		Treat	Treated	
	Untreated	nonwastewa	wastewater .	
BDAT List	<u>waste</u>	Total	TCLP	(scrubber water
Constituent	(ppm)	(ppm)	(mg/l)	(mg/l)
<u>Metals</u>				
Antimony	<17	<17	0 040	0 040
Arsenic	2.6	11	<0.020	1 6
Barıum	150	72	0.41	1 1
Beryllium	< 0 5	< 0 5	< 0 005	0 001
Cadmium	3.5	<2 0	<0.020	0 46
Chromium	8 6	7 7	<0 055	0 89
Copper	38	8€	0 070	0 35
Mercury	0 64	<1.25	< 0 002	5 0 54
Nickel	4 5	3 9	0.020	0 54
Lead	190	<21	< 0 21	J.9
Selenium	1.1	2 8	<0.020	0 021
Silver	<3 5	< 3 5	< 0 035	0 0202
Thallium	3 3	<2.5	< 0 025	4 0
Vanadium	1.9	4.8	<0.040	0 060
Zinc	200	2 5	0 002	ъ 2

Reference Onsite Engineering Report for K001-Creosote

Table 3-7 Rotary Kiln Incineration of K001 - PCP Sample Set No 7

BDAT List Constituent	Untreated waste (ppb)	Treated nonwastewater Total (ppb)	Treated wastewater (ug/1)
Volatile Organics			
To luene	16	<10	<10
	Treated		
	Untreated	nonwastewater	Treated
	waste	Total	<u>wastewater</u>
	waste (ppm)	Total (ppm)	wastewater (ug/l)
Sem.volatile Organic Acenaphthene Anthracene Benz(a)anthracene Benzo(a)pyrene	(ppm)		
Acenaphthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k)	(ppm) 13.000 9,300 <2,500 <250	<2.5 <2.5 <2.5 <2.5 <2.5	<50 <50 <10 <50
Acenaphthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluoroanthrene	(ppm) 13.000 9,300 <2,500 <250	<2.5 <2.5 <2.5 <2.5 <2.5 <2.5	<50 <50 <10 <50
Acenaphthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluoroanthrene Chrysene	(ppm) 15.000 9,300 <2,500 <250 940 <2,500	<2.5 <2.5 <2.5 <2.5 <2.5 <2.5	<50 <50 <10 <50 <50 <10 <50 <50 <50 <50 <50 <50
Acenaphthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluoroanthrene Chrvsene Fluoranthrene	(ppm) 13.000 9,300 <2,500 <250 940 <2,500 13,000	<2.5 <2.5 <2.5 <2.5 <2.5 <2.5 <2.5	<50 <50 <10 <50 <50 <50 <50 <50 <50 <50 <50 <50
Acenaphthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluoroanthrene Chrysene Fluoranthrene Fluorene	(ppm) 13.000 9.300 <2.500 <250 940 <2.500 13.000 8.200	<2.5 <2.5 <2.5 <2.5 <2.5 <2.5	<50 <50 <10 <50 <50 <10 <50 <50 <50 <50 <50 <50
Acenaphthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluoroanthrene Chrvsene Fluoranthrene	(ppm) 13.000 9,300 <2,500 <250 940 <2,500 13,000	<pre></pre>	<pre></pre>
Acenaphthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluoroanthrene Chrvsene Fluoranthrene Fluorene Naphthalene	(ppm) 13.000 9,300 <2,500 <250 940 <2,500 13.000 8,200 26,000	<pre></pre>	<50 <50 <10 <50 <50 <50 <50 <50 <50 <50 <50 <50 <5

Table 3-7 (continued)

Sample Set No. 7

	Untreated	Trea _nonwaste		Treated	
BDAT List	waste	Total	TCLP	wastewater	
Constituent	(ppm)	(ppm)	(mg/1)	(mg/l)	
<u>Metals</u>					
Ant imony	<30	<30	<0.3	<0.3	
Arsenic	2 9	0 8	<0.015	<0.01	
Barıum	30	74	0.32	0 12	
Beryllium	< 0 5	< 0 5	<0.005	< 0 005	
Cadmium	0.5	<15	< 0 015	<0 015	
Chromium	1.5	8 2	< 0 045	< 0 045	
Copper	6.7	6 δ	<0.05	0 15	
Lead	7.8	5 2	0.021	0 021	
Mercury	0.11	<0.001	<0.001	<0.001	
Nickel	< 10	< 10	<0.1	< 0 1	
Selenium	<2 5	<2 5	<0.025	< 0 025	
Silver	<4.5	<4 5	<0.045	< 0 045	
Thallium	<1.5	<1 5	<0.05	< 0 015	
Vanadium	<10	< 10	<0.1	<0 1	
Zinc	64	11	<0 03	1.1	

Reference: Onsite Engineering Report for K001-Pentachlorophenol

Table 3-8 Rotary Kiln Incineration of K001 - PCP Sample Set No 8

BDAT List Constituent	Untreated <u>waste</u> (ppb)	Treated nonwastewater Total (ppb)	Treated wastewater (ug/l)
Volatile Organics			
To luene	10	<10	<10
	llan on the d	Treated	Tuestad
	Untreated waste	<u>nonwastewater</u> Total	Treated w <u>aste</u> water
	- Marie 2011		
·-	(ppm)	(ppm)	(ug/1)
Semivolatile Organic Acenapthene Anthracene Benz(a)anthracene	\$ 18,000 13,000 3,400	<2 5 <2 5 <2 5	<50 <50
Acenapthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k)	\$ 18,000 13,000 3,400 940	<2 5 <2 5 <2 5 <2 5	<50 <50 <10 <50
Acenapthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluroanthrene	\$\frac{1}{2},000\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	<2 5 <2 5 <2 5 <2 5 <2 5 <2 5	<50 <50 <10 <50
Acenapthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluroanthrene Chrysene	18,000 13,000 3,400 940 2,300 ,600	<2 5 <2 5 <2 5 <2 5 <2 5 <2 5 <2 5	<50 <50 <10 <50 <50
Acenapthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluroanthrene Chrysene Fluoranthrene	\$\frac{1}{2},000 13,000 13,000 3,400 940 2,300 1,600 21,000	<2 5 <2 5 <2 5 <2 5 <2 5 <2 5 <2 5 <2 5	<50 <50 <10 <50 <50 <50 <50
Acenapthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluroanthrene Chrysene Fluoranthrene Fluorene	\$\frac{5}{18,000}\$ 18,000 13,000 3,400 940 2,300 .,600 21,000 12,000	<2 5 <2 5 <2 5 <2 5 <2 5 <2 5 <2 5 <2 5	<50 <50 <10 <50 <50 <50 <50
Acenapthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluroanthrene Chrysene Fluoranthrene Fluorene Naphthälene	\$\frac{1}{2},000 13,000 13,000 3,400 940 2,300 .600 21,000 12,000 43,000	<2 5 <2 5 <2 5 <2 5 <2 5 <2 5 <2 5 <2 5	<50 <50 <10 <50 <50 <50 <50 <50
Acenapthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluroanthrene Chrysene Fluoranthrene Fluorene	\$\frac{5}{18,000}\$ 18,000 13,000 3,400 940 2,300 .,600 21,000 12,000	<2 5 <2 5 <2 5 <2 5 <2 5 <2 5 <2 5 <2 5	<50 <50 <10 <50 <50 <50 <50

Table 3-8 (continued)

Sample Set No 8

		Trea	ted			
•	Untreated	nonwast	ewater	Treated		
BDAT List	<u>waste</u>	Total	TCLP	wastewater		
Constituent	(ppm)	(ppm)	(mg/1)	(mg/1)		
<u>Metals</u>						
Ant imony	<30	<30	<0.3	< 0 3		
Arsenic	2.3	0.6	<0.015	0 12		
Barıum	19	21	0.19	0 24		
Beryllium	< 0 5	< 0 5	<0.005	< 0 005		
Cadmium	0 6	< 1 5	<0.015	< 0 015		
Chromium	2 7	1 1	<0.045	< 0 045		
Copper	11	3.0	<0.05	0 09		
Lead	. 11	1 2	< 0 01	0 18		
Mercury	0 16	< 0 001	< 0 001	< 0 001		
Nickel	<10	<10	< 0.1	< 0 1		
Selenium	<2 5	<2 5	<0.025	< 0 025		
Silver	<4 5	< 4 5	< 0 045	< 0 045		
Thallium	<1.5	<1 5	< 0 05	< 0 015		
Vanadium	<10	<10	<0.1	< 0 1		
Zinc	58	2.1	<0.03	0 61		

Reference: Onsite Engineering Report for K001-Pentachlorophenol

Table 3-9 Rotary Kiln Incineration of K001 - PCP

Sample Set No 9

BDAT List Constituent	Untreated waste (ppb)	Treated nonwastewater Total (ppb)	Treated <u>wastewater</u> (ug/l)
Volatile Organics			
To luene	39	<10	<10
	Untreated waste	Treated <u>nonwastewater</u> Total	Treated wastewater
	(ppm)	(ppm)	(ug/1)
Semivolatile Organic		(ppm)	(ug/l)
Semivolatile Organic Acenapthene		(ppm) 	(ug/1)
	<u>s</u>		
Acenapthene	<u>s</u> 1+,000	<2.5	<50
Acenapthene Anthracene	<u>s</u> 14,000 5,500	<2.5 <2.5	<50 <50
Acenapthene Anthracene Benz(a)anthracene Benzo(a)pyrene	14,000 5,500 2,500	<2.5 <2 5 <2 5	<50 <50 <10
Acenapthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k)	14,000 5,500 2,500 620	<2.5 <2.5 <2.5 <2.5	<50 <50 <10 <50
Acenapthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluroanthrene	14,000 5,500 2,500 620	<2.5 <2.5 <2.5 <2.5 <2.5	<50 <50 <10 <50
Acenapthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluroanthrene Chrysene	14,000 5,500 2,500 620 1,600 42,500	<2.5 <2.5 <2.5 <2.5 <2.5 <2.5	<50 <50 <10 <50 <50
Acenapthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluroanthrene Chrysene Fluoranthrene	14,000 5,500 2,500 620 1,600 <0,500 15,000	<2.5 <2 5 <2 5 <2.5 <2.5 <2.5 <2.5	<50 <50 <10 <50 <50 <50 <50
Acenapthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluroanthrene Chrysene Fluoranthrene Fluorene	14,000 5,500 2,500 620 1,600 -2,500 15,000 9,000	<2.5 <2 5 <2 5 <2.5 <2.5 <2.5 <2.5 <2 5 <2 5 <2 5	<50 <50 <50 <50 <50 <50 <50 <50
Acenapthene Anthracene Benz(a)anthracene Benzo(a)pyrene Benzo(b &/or k) fluroanthrene Chrysene Fluoranthrene Fluorene Naphthalene	14,000 5,500 2,500 620 1,600 +2,500 15,000 9,000 37,000	<2.5 <2 5 <2 5 <2.5 <2.5 <2.5 <2 5 <2 5 <2 5 <2 5 <2 5 <2 5	<50 <50 <10 <50 <50 <50 <50

Table 3-9 (continued)

Sample Set No. 9

		Treat	ted	
	Untreated	nonwast	ewater	Treated
BDAT List	<u>waste</u>	Total	TCLP	wastewater
Constituent	(ppm)	(ppm)	(mg/1)	(mg/1)
<u>Metals</u>				
Ant imony	<30	<30	<0.3	<0.3
Arsenic	1.1	0 4	<0.015	0 11
Barıum	17	21	0.25	0 39
Beryllium	<0.5	< 0 5	<0.005	< 0 005
Cadmium	0.4	<1 5	<0.015	<0.015
Chromium	2 1	1 2	<0.045	0.045
Copper	10	2	< 0 05	0 07
Lead	6.3	0 96	<0.01	0 20
Mercury	0.064	<0.001	<0.001	0.003
Nicke?	<10	<10	<0.1	<0.1
Se len ium	<2 5	<2.5	< 0 025	< 0 025
Silver	< 4 5	< 4 5	<0.045	< 0 045
Thallium	<1.5	<1.5	<0.05	< 0.015
Vanadıum	<10	<10	<0.1	< 0.1
Zinc	30	2.1	<0.03	0.88

Reference Onsite Engineering Report for K001-Pentachlorophenol.

Table 3-10 Incinerator Operating Data for KOO1 - Creosote Sample Set Number 1 and 2

		K001-creosote feed rate, Temperature, *F			Kilm rotation	Stack gas concentrations			
Parameter	Sample collected	1b/h	Kıln	Afterburner	speed, rpm	02, 7	CO, ppm	co ₂ . 5	
0/5/87									
nln shakedown (before K001-C feed) 0/6/87	Scrubber pretest water	0	1720-1917	1984-2019	0 25				
Start feed 11:03		180			0.25				
Begin test 12:10	Feed (a)	180	1694-1873	1932-2016	0.25	4-9	12-27	8-12	
12:30		180	1771-1886	2008-2031	0.25	4-12	22-32	3-11	
13.00		180	1828-1863	2014-2031	0.25	4-20	18-33	5-12	
13:30		180	1835-1874	2014-2035	0 25	5-8	30-34	1-11	
13 40	Ash	180	1835-1838	2030-2033	0.25	6-8	32-34	9-10	
	Recycle (b)								
14:00	Feed (a)	180	1790-1860	1996-2033	0.25	5-9	32-37	8-11	
14:30		180	1800-1913	1996-2034	0.25	5-8	32-37	9-11	
15.00		180	1838-1906	2019-2035	0.25	6-8	34-37	8-11	
15:25	Ash	180	1795-1838	2006-2019	0.25	7-9	30-37	8-10	
	Recycle (b)								

⁽a) The feed samples were collected while the fiber packs were being packed. Six drums were used for the two days of testing. K001-C waste from two drums was incinerated for each sample set.

Reference Onsite Engineering Report for K001-Creosote

⁽b) The scrubber recycle tank was not blown down during the test series. Makeup water was added to replace water loss through the exhaust system. No chemicals were added to the scrubber water system. The recycle water samples were taken from the scrubber recirculation line.

Table 3-11 Incinerator Operating Data for KOO1 - Creosote Sample Set Number 3 and 4

Parameter Sample collected		K001-creosote feed rate,				Stack gas concentrations			
	lb/h	Kiln	Afterburner	rotation speed, rpm	02, 1	CO, ppm	co ₂ , /		
10/6/87									
Continued feed									
17 10	Feed (a)	180			0 25				
17 30		180	1775-1840	1985-2002	0 25	6-10	39-42	8-10	
18:00		180	1827-1875	2002-2026	0 25	6-9	31-42	8-11	
18.30		180	1827-1875	2007-2026	0 25	6-9	39-41	8-10	
18:40	Ash	180	1822-1871	1993-2009	0.25	7-9	39-41	8-9	
	Recycle (b)								
19:00	Feed (a)	180	1819-1826	1993-1997	0 25	7-9	37-41	8-10	
19.30		180	1825-1878	1997-2008	0 25	7-9	31-42	8-11	
2 .00	O Recycle (b)	180	1859-2028	1995-2093	0 25	4 - 9	39-720	8-13	
20:15	Ash	180	1824-1859	1986-1995	0.25	7-10	35-38	8-10	

- (a) The feed samples were collected while the fiber packs were being packed. Six drums were used for the two days of testing. K001-C waste from two drums was incinerated for each sample set.
- (b) The scrubber recycle tank was not blown down during the test series Makeup water was added to replace water loss through the exhaust system. No chemicals were added to the scrubber water system. The recycle water samples were taken from the scrubber recirculation line.

Reference: Onsite Engineering Report for K001-Creosote.

		K001-creosote feed rate,	Temperatu	re. "F	Kiln rotation	Stack gas concentrations			
Parameter Sample collected	lb/h	Kiln	Afterburner	speed, rpm	02, %	СО, ррт	co ₂ , %		
0/7/87									
Start feed 11.00		180			0 25				
Begin test 11 30	Feed (a)	180	180€ 1841	2012 - 2020	0 25	4 - 8	15-18	9-12	
12.00		180	1823-1912	2017-2039	0.25	2-9	15-30	9-13	
12.30	Ash	180	1830-1886	2019-2025	0 25	4-8	17-24	9-11	
	Recycle (b)								
13:00	Feed (a)	180	1845-1910	2011-2025	0.25	5-9	19-25	9-10	
13:30		180	1829-1845	1999-2011	0 25	6-9	18-22	9-10	
14.00		180	1827-1838	1997-2009	0 25	7-9	18-22	9-10	
14 30		180	1827-1857	1999-2022	0 25	7-9	15-23	9-10	
14 45	Ash	180	1844-1899	2022-2034	0 25	5-8	22-25	9-10	
	Recycle (b)								
15 00		180	1896-1900	2026-2033	0 25	5-9	22-25	9-11	

⁽a) The feed samples were collected while the fiber packs were being packed. Six drums were used for the two days of testing. K001-C waste from two drums was incinerated for each sample set.

Reference: Onsite Engineering Report for KOO1-Creosote.

⁽b) The scrubber recycle tank was not blown down during the test series. Makeup water was added to replace water loss through the exhaust system. No chemicals were added to the scrubber water system. The recycle water samples were taken from the scrubber recirculation line.

Table 3-13 Incinerator Operating Data for K001-PCP Sample Sets #7 - #9

Sample Set #7

	Sample	Tempe	erature, 4 1F	Kiln rotation	Stack gas concentrations			
Parameter	collected	Kiln	Afterburner	speed, rpm	02, %	CO, p	pm CO ₂ , %	
6/26/87								
Start feed (86 lb/h) Feed (a)							
drum #1 15 30	Makeup water							
15 45								
16 00		1650-1682	1840-1869	0.2	5-9	< 1	4->10	
16 15		1720-1803	1823-1866	0.2	5-8	<1	7->10	
16 30		1767-1826	18 69-1875	0.2	5-9	< 1	6->10	
16 45		1777-1820	1872-1879	0.2	6-8	<1	8-10	
17 00		1801-1873	1876-1895	0.2	5-8	< 1	9->10	
17 15		1813-1899	1889-1903	0.2	4-9	<1	8->10	
17 30		1793-1936	1890-1914	0 2	4-9	<1	8->10	
17 45		1861-1914	1885-1912	0 2	6-8	<1	9->10	
18 00	•	1823-1914	1868-1888 .	0 2	5-16	<1	9->10	
18 15		1782-1892	1859-1871	0 2	4-16	< 1	9->10	
18 3 0 (E)	1877	1839	0.2	3-8	< 1	8->10	
18 45 (b		1924	1846	0.2	3-8	<1	8->10	
19 00 (b		1970	1857	0.2	3-9	<1	8->10	
19 15 (E		2046	2033	0.2	3-10	<1	8->10	

⁽a) The feed samples were collected while the fiber packs were being packed. Sample number CKO1P-1-AX represents feed to the kiln during sample set 7, sample number CKO1P-2-AX represents feed to the kiln during sample set 8, and sample number CKO1P-3-AX represents feed to the kiln during sample set 9

Reference Onsite Engineering Report for K001-Pentachlorophenol.

⁽b) Only one temperature value was available during this time period.

⁽c) The ash samples were collected from the ash bin after the ash cooled.

Table 3-14 Performance Data for Chemical Precipitation and Filtration on Mixed Waste Sampled by EPA

				Concentrat				
	Sample	Set #1	Sample	Set #2	Sample	Set #3	Sample	Set #4
	Treatment	Treatment			Treatment		Treatment	
Constituent/parameter	tank composite	Filtrate	tank composite	Filtrate	tank composite	Filtrate	tank composite	Filtrate
BDAT Metals	•							
Ant imony	<10	<1	< 10	< 1	<10	< 1	<10	+
Arsenic	<1	< 0 1	<1	< 0 1	<1	<0.1	<1	<1
Barıum	<10	< 1	< 10	<1	<10	3 5	< 10	< 10
Beryllıum	<2	<0.2	<2	< 0 2	<2	< 0 2	<2	<2
Cadmium	13	<0.5	10	< 0.5	<5	<0.5	< 5	<5
Chromium (hexavalent)	893	0.011	807	0.190	775	_a	0 6	0 042
Chromium (total)	2,581	0 12	2,279	0 12	1,990	0 20	556	0 10
Copper	138	0 21	133	0 15	133	0 21	88	0 07
.ead	64	< 0 01	54	< 0 01	< 10	< 0 01	< 10	< 0 01
Mercury	< 1	< 0.1	<1	< 0 . 1	<1	< 0 1	<1	<1
lickel	471	0.33	470	0 33	16,330	0.33	6,610	0 33
Selenium	<10	< 1	<10	< 1	<10	< 1	<10	<10
Silver	<2	< 0 2	2	< 0 2	< 2	< 0 3	<2	<2
[hallıum	< 10	<1	< 10	< 1	< 10	<1	<10	< 10
Zinc	116	0 125	4	0 115	3 9	0 140	84	1 62
Other Parameters								
Total organic carbon	2700		2800		500		2900	
Total solids	-		-		-		-	
Total chlorides	-		-		-		-	
Total organic halides	2500		3600		0		900	

	Samo le	Set #5	Sample	Set #6	Sample	Set #7	Sample	Set #8
	Treatment		Treatment		Treatment		Treatment	
Constituent/parameter	tank composite	Filtrate						
BDAT Metals								
Ant imony	<10	< 1	< 10	< 1	<10	< 1	<10	<1
Arsenic	<1	< 0 1	<1	< 0.1	<1	< 0 1	<1	<0.1
Barium	<10	< 1	<10	<2	<10	< 1	<10	< 1
Beryllium	<2	< 0 2	<2	< 0 2	<2	< 0.2	<2	< 0 2
Cadmium	<5	<0.5	< 5	<0.5	10	<0.5	<5	< 0 5
Chromium (hexavalent)	917	0 058	734	_a	769	0.121	0.13	<0.01
Chromium (total)	2,236	0.11	2,548	0.10	2,314	0.12	831	0 15
Copper	91	0 14	149	0 12	72	0.16	217	0 16
Lead	18	< 0 01	< 10	< 0 01	108	< 0 01	212	<0.01
Mercury	1	< 0 1	<1	< 0 1	<1	< 0 01	<1	< 0 1
Nickel	1,414	0.310	588	0 33	42 6	0.40	669	0 36
Selenium	<10	< 1	<10	< 1	<10	< 1	<10	< 1
Silver	<2	<0 2	<2	< 0 2	<2	<0.2	<2	<0.2
Thallium	<10	< 1	<10	< 1	< 10	<1	<10	<1
Zinc	71	0 125	4	0 095	171	0 115	151	0 130
Other Parameters								
Total organic carbon	200		700		3400		5900	
Total solids	-		-		-		-	
Total chlorides	-		-		-		-	
Total organic halides	0		700		1900		800	

Table 3-14 (Continued)

	Concentration (ppm)								
	Sample	Set #9	Samp le	Set #10		Set #11			
	Treatment		Treatment		Treatment				
Constituent/parameter	tank composite	Filtrate	tank composite	Filtrate	tank composite	Filtrate			
BDAT Metals									
Ant imony	<10	<1	<10	<1	<10	<1.00			
Arsenic	<1	< 0 1	<1	<0.1	<1	< 0.10			
Barıum	< 10	< 1	<10	<1	<12	<1.00			
Beryllium	<2	< 0 2	<2	<0.2	<2	< 0 20			
Cadmium	<5	< 0 5	<5	<0.5	23	< 5			
Chromium (hexavalent)	0.07	0.041	0.08	0.106	0.30	< 0 01			
Chromium (total)	939	0.10	395	0.12	617	0.18			
Copper	225	0.08	191	0 14	137	0.24			
Lead	<10	<0.01	<10	<0.01	136	<0.01			
Mercury	<1	< 0.1	<1	< 0 1	<1	< 0 10			
Nickel	940	0 33	712	0.33	382	0 39			
Selenium	<10	<1.0	<10	< 1	<10	<1.00			
Silver	<2	<0.2	<2	< 0 2	<2	< 0 2			
Thallium	<10	<1.0	<10	< 1	<10	<1.00			
Zinc	5	0.06	5	0.070	135	0.100		•	
Other Parameters									
Total organic carbon	2100		0		52				
Total solids	-		_		-				
Total chlorides	-		_		-				
Total organic halides	0		<300		300				

^aHexavalent chromium was actually treated by chromium reduction prior to chemical precipitation and sludge filtration - = Not analyzed.

Note. Design and Operating Parameters are as follows.

pH during chromium reduction - 8.5 to 9.0.

Reducing agent - ferrous iron.

Ratio of reducing agent to hexavalent chromium - 3.2 to 10.

pH during chemical precipitation - 8 to 10.

Precipitation agents - lime and sulfide.

Filter type - vacuum filter.

Reference: Onsite Engineering Report for Envirite

Table 3-15 EPA Collected Total Composition Data for Untreated FOO6 Waste

Consti	tuent		Cor	ncentration in	Raw Waste Samp	ole - F006 (ppm)			
	#1	#2	#3	#4	#5	#6	#7	#8	#9
Barıum			85 5		14.3		-+	15.3	19
Cadmium	~ ~	31.3	67.3	1 31	720	7.28	5.39	5.81	
Chromium		755	716	* *	12200	3100	42900		
Copper		7030	693	1510	160	1220	10600	17600	27,400
_ead		409	257	88 5	52	113	156	1.69	24,500
lickel	435	989	259	374	701	19400	13000	23700	5730
Silver		6 62	38 9	9 05	5 28	4.08	12 5	8.11	
Zinc	1560	4020	631	90200	35900	27800	120	15700	322

^{1 -} Wastewater treatment sludge cake - no free liquid

Reference: CWM Technical Note 87-117.

^{2 -} Site closure excavation mud at auto part manufacturer. The waste sample is a mixture of F006 and F007.

^{3 -} Waste treatment sludge from aircraft overhaul facility. The waste sample is a mixture of F006, D006, D007, and D008.

^{4 -} Zinc electroplating sludge.

^{5 -} Filter cake from electroplating wastewater treatment

^{6 -} Sludge from treatment of Cr. Cu, Ni, and Zn plating

^{7 -} Wastewater treatment sludge from plating on plastics

^{8 -} Wastewater treatment sludge

^{9 -} To be provided

Table 3-16 EPA Collected TCLP Data for Untreated F006 Waste $^{\rm a}$

Const ₁	tuent				TCLP Concer	ntration (ppm)		
	#1	#2	#3	#4	#5	#6 .	#7	#8	#9
Barium			1 41		0 38			0 53	0 28
Cadmium		2.21	1 13	0.02	23.6	0.03	0.06	0 18	
Chromium		0 76	0.43		25 3	38.7	360	- -	
Copper		368		4.62	1 14	31 7	8 69	483	16 9
ead		10 7	2 26	0 45	0 45	3 37	1 0	4 22	50 2
Vickel	0.71	22 7	1.1	0 52	9 78	730	152	644	16 1
Silver		0.14	0.20	0 16	0 08	0.12	0.05	0 31	
inc	0 16	219	5 41	2030	867	1200	0.62	650	1 29

 $^{^{\}mathrm{a}}\mathrm{See}$ Table 3-15 for sample descriptions.

Reference CWM Technical Note 87-117

Table 3-17 EPA Collected TCLP Data for F006 Stabilized $Residues^a$

Consti	tuent				Concer	ntration (ppr	n)		
	#1	#2	#3	#4	#5	#6	#7	#8	#9
Mix ratio	0.2	0.5	0.2	1 0	0.5	0.5	0.5	0.5	0.5
Barıum			0.33		0.23			0 27	0.08
Cadmium		0.01	0.06	0.01	0.01	0.01	0.01	0.01	
Chromium		0.39	0 08		0.03	0.38	1.21		
Copper		0.25		0 15	0.27	0 29	0.42	0.32	0 46
Lead		0.36	0 30	0 21	0 34	0.36	0.38	0.37	0 27
Nickel	0 04	0.03	0 23	0 02	0.03	0.04	0.10	0.04	0 02
Silver		0.05	0 20	0 03	0.04	0.06	0.05	0.05	
Zanc	0 03	0 01	0.05	0 01	0.04	0.03	0.02	0.02	<0.01

 $^{^{\}rm a}{\rm See}$ Table 3-15 for sample descriptions of all of the samples of raw waste.

Reference CWM Technical Report 87-117

4. IDENTIFICATION OF BEST DEMONSTRATED AVAILABLE TECHNOLOGY (BDAT) FOR KOO1

This section presents the Agency's methodology for identifying the best demonstrated available technology (BDAT) for treatment of K001 based on the performance data presented in Section 3. The demonstrated technologies for treatment of BDAT list organic constituents present in K001 wastes are incineration and fuel substitution. For BDAT list metals in nonwastewater forms such as scrubber water treatment nonwastewater residuals and incinerator ash, the demonstrated treatment technology is stabilization. Chemical precipitation and filtration is the demonstrated treatment train for BDAT list metals in wastewater forms of K001 such as scrubber water from incineration.

As stated in the Introduction, BDAT is selected based on treatment performance data available to the Agency. Prior to being used to establish treatment standards, performance data are screened to determine whether they meet the requirements of the BDAT program. First, the design and operating data collected for each data set are examined, and data points or data sets that reflect a poorly designed treatment system, or a system that was not well operated at the time of data collection, are not used in the development of treatment standards. In addition, data are screened with regard to the quality assurance/quality control measures (QA/QC) and whether the appropriate analytical methods were used to assess the performance of the treatment technology. All remaining performance data are then adjusted based on analytical recovery

values, which, in turn are based on laboratory quality assurance/quality control analyses in order to take into account analytical interferences associated with the chemical makeup of the sample. Finally, in cases where the Agency has performance data on treatment of a listed waste using more than one demonstrated technology, the treatment values are compared by the analysis of variance test (ANOVA), as presented in Appendix A. This test will determine if one technology performs significantly better than another. This was not the case with KOO1, since data from only one treatment technology were available.

4.1 Review of Performance Data

The available treatment data described in Section 3.0 were reviewed and assessed with regard to the design and operation of the treatment systems, the analytical testing, and the quality assurance/quality control analyses of the data. In general, all of the performance data collected for rotary kiln incineration of K001 were of sufficient quality to develop treatment standards. Design and operating data were collected for the rotary kiln incineration systems used for destroying BDAT list organic constituents in K001 wastes. These data indicate that the systems were well designed and well operated during the test burns. In addition, the proper analytical tests were performed for the untreated wastes and the treated residuals. Specifically, because incineration is a destruction technology for organics, total constituent concentration of organics is used to measure treatment performance.

In one instance, the analytical quality assurance/quality control data collected during the analyses of the KOOl incineration samples were not of sufficient quality for use in developing BDAT treatment standards. Specifically, the matrix spike recoveries for pentachlorophenol in the scrubber water from the KOOl-PCP test burn were below acceptable limits (20%), and therefore, could not be used to develop treatment standards. In this case, recovery data for pentachlorophenol in the scrubber water were transferred from the most appropriate source, KOOl-creosote. Section 6 of this document presents the recovery data used in each case and Appendix B contains all recovery data developed for KOOl waste.

The performance data transferred from stabilization testing of F006 contained the required data on design and operation, QA/QC, and the proper analytical testing (total composition and TCLP for the untreated waste and TCLP for the treated waste). These treatment data were used in the development of treatment standards for K001. None of the eleven data sets for treatment of the wastewaters by chemical precipitation and filtration were deleted. Design and operating data collected during the sampling of this treatment system did not indicate that the system was poorly designed or operated. In addition, analyses were performed for total composition in the untreated wastes as well as the treated waste. However, matrix spike recovery data were not available for the BDAT list metal constituents in these waste streams. Matrix spike recovery data

were transferred from the Onsite Engineering Report for Horsehead Resource Development Co. for K061. The recovery data from the TCLP extracts of the treated K061 residuals were used because the waste matrices were determined to be similar.

4.2 Accuracy Correction of Performance Data

After the screening tests, EPA adjusted the data values based on the analytical recovery values in order to take into account analytical interferences associated with the chemical makeup of the treated sample. In developing recovery data (also referred to as accuracy data), EPA first analyzed the sample for a constituent and then added a known amount of the same constituent (i.e., spike) to the waste material. The total amount recovered after spiking minus the initial concentration in the sample divided by the amount added is the recovery value.

In general, a matrix spike recovery is determined from the result of one matrix spike performed for each individual constituent. Such is the case for BDAT list metals and selected BDAT list volatile and semivolatile constituents. However, for constituents for which no matrix spike recovery was performed, the recovery data were determined from the average matrix spike recoveries of the appropriate group of constituents for which recovery data were available. For example, no matrix spike was performed for xylenes; the matrix spike recovery data used for xylenes were the result obtained by averaging the matrix spike recoveries for all BDAT list volatile constituents that had recovery data.

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Where matrix spikes were not performed for a BDAT list semivolatile constituent, a matrix spike recovery for that constituent was calculated based on semivolatile constituents for which there were recovery data for two matrix spikes. The lower of the two average matrix spike recoveries of semivolatile constituents was used in this case. For example, no matrix spike recovery was performed for naphthalene, a base/neutral fraction semivolatile. The recovery data for naphthalene were developed after averaging the matrix spike recoveries calculated for all base/neutral fraction semivolatiles in both the first matrix spike and the duplicate spike. The lower average matrix spike recovery was selected to calculate the correction factor for naphthalene.

The accuracy correction factors are calculated from the recovery data. In general, the reciprocal of the lower recovery value, divided by 100, yields the correction factor. The accuracy corrected values are obtained by multiplying the uncorrected data value by the correction factor. These adjusted values were then used to calculate treatment standards for BDAT list constituents as presented in Section 6.

Appendix B presents the analytical methods and quality assurance/quality control data used to develop the recovery values for each constituent.

4.3 BDAT for Treatment of Organics

After analyzing the accuracy corrected data, EPA has determined that rotary kiln incineration achieves a level of performance that represents organic treatment by BDAT for KOO1. EPA would not expect the level of

performance to be improved by other forms of incineration such as fluidized bed or fixed hearth systems because rotary kiln incineration destroyed BDAT list organics to concentrations below their detection limits. In addition, EPA believes that well-designed and well-operated fuel substitution systems could not achieve better treatment since they operate at approximately the same temperatures and turbulent conditions as an incineration system.

In addition to being the "best demonstrated" technology for BDAT list organics in KOO1, rotary kiln incineration is also "available" because it is commercially available or can be purchased from a proprietor, and it provides substantial reduction of the concentration of the BDAT list organics. Because EPA has determined that rotary kiln incineration is "best," "demonstrated," and "available," it is the technology basis for treatment standards for BDAT list organic constituents present in KOO1 wastes.

4.4 BDAT for Treatment of Metals

EPA has determined that stabilization achieves a level of performance that represents treatment by BDAT for BDAT list metals in nonwastewater KOO1 treatment residuals such as wastewater treatment nonwastewater residuals and incinerator ash. The Agency has no reason to expect that the level of performance could be improved, since stabilization is the only demonstrated treatment technology identified by EPA. For BDAT list metals in the wastewater residual, only one technology treatment train

was identified as being demonstrated (i.e., precipitation and filtration). For BDAT list metals in scrubber water, therefore, this treatment train is designated as BDAT. The Agency has no reason to expect that the level of performance for BDAT list metals in wastewaters could be improved beyond the specified BDAT treatment level.

Stabilization of the nonwastewater treatment residuals and filtration of the wastewater are judged to be available to treat BDAT list metals present in K001 treatment residues because the treatments are commercially available or can be purchased from a proprietor and these treatments provide substantial reduction of the concentration and/or leachability of hazardous metal constituents. As a result, stabilization of K001 nonwastewater residuals and chemical precipitation and filtration for K001 wastewaters are the technology basis for BDAT list metals in K001 wastes.

SELECTION OF REGULATED CONSTITUENTS

This section presents the methodology and rationale for selection of the constituents that are being proposed for regulation in wastewater and nonwastewater forms of KOOl wastes.

As discussed in Section 1, the Agency has developed a list of hazardous constituents (Table 1-1) from which the pollutants to be regulated are selected. The list is a "growing list" that does not preclude the addition of new constituents as additional key data and information parameters become available. The list is divided into the following categories: volatile organics, semivolatile organics, metals, inorganics other than metals, organochlorine pesticides, phenoxyacetic acid herbicides, organophosphorus pesticides, PCBs, and dioxins and furans. Also discussed in Section 1 is EPA's process for selecting constituents to regulate. In general, this process consists of identifying constituents in the untreated waste that are present at treatable concentrations and then regulating the constituents in that group necessary to ensure effective treatment. Below is a discussion that details how EPA arrived at the list of constituents to be regulated for K001.

5.1 BDAT List Constituents Detected in Untreated KOO1 Waste

Of the 232 constituents on the BDAT list, 31 were detected in the untreated K001 waste. Table 5-1 shows the specific constituents that were analyzed and detected in the untreated K001, as well as the detection limits. For the constituents not detected, it was assumed that

Table 5-1 BDAT List Constituents in Untreated K001 Waste*

BDAT			D = Detected	
reference			ND = Not detected	Detection
no	Parameter	Units	NA = Not analyzed	limit
	<u>Volatiles</u>			
222	Acetone	ppb	ND	250
1.	Acetonitrile	ppb	ND	1000
2.	Acrolein	ppb	ND	2500
3	Acrylonitrile	ppb	ND	50
4	Benzene	ppb	D	50
5.	Bromodichloromethane	ppo	ИД	50
6	Bromomethane	ppb	ND	50
223	n-Butyl alcohol	ppb	NA	-
7	Carbon tetrachloride	ppb	ND	50
8.	Carbon disulfide	ppb	ND	50
9	Chlorobenzene	ppb	ND	50
10	2-Chloro-1,3-butadiene	ppb	ND	0 25
11.	Chlorodibromomethane	ppb	ND	50
12.	Chloroethane	ppb	ND	50
13	2-Chloroethyl vinyl ether	agg	ND	500
14	Chloroform	ppb	ND	50
15	Chloromethane	ppb	ND	50
16.	3-Chloropropene	ppb	ND	50
17.	1,2-Dibromo-3-chloropropane	aqq	ND	50
18.	1,2-Dibromoethane	ppb	ND	50
19	Dibromomethane	aqq	ND	50
20.	Trans-1,4-Dichloro-2-butene	ppb	ND	50
21	Dichlorodifluoromethane	ppb	ND	50
22.	1,1-Dichloroethane	dad	ND	50
23.	1,2-Dichloroethane	ppb	ND	50
24	1,1-Dichloroethylene	ppb	ND	50
25	Trans-1,2-Dichloroethene	ppb	ND	50
26	1,2-Dichloropropane	ppb	ND	50
27	Trans-1,3-Dichloropropene	ppb	ND	50
28.	cis-1,3-Dichloropropene	ppb	ND	50
29.	1,4-Dioxane	ppb	ND	250
224	2-Ethoxyethanol	ppb	NA	-
225	Ethyl acetate	ppb	ND	50
226.	Ethyl benzene	ppb	D	10
30.	Ethyl cyanide	ppb	ND	0.5
227	Ethyl ether	ppb	ND	250
31	Ethyl methacrylate	ppb	ND	250
214	Ethylene oxide	ppb	ND	250
32.	Iodomethane	ppb	ND	100

Untreated KOOl was sampled on two occasions and originally presented in two Onsite Engineering Reports (OER). These data reflect the constituents detected in any of the samples while the detection limit listed is the highest of the two presented in the OERs for KOOl.

⁻ No detection limit available.

Table 5-1 (continued)

BDAT			D = Detected	
refe re nce			ND = Not detected	Detection
no ·	Parameter	Units	NA = Not analyzed	limit
	<u>Volatiles</u> (continued)			
33	Isobutyl alcohol	ppb	ND	1
228.	Methano 1	dqq	NA	-
34	Methyl ethyl ketone	ppb	ND	250
229	Methyl isobutyl ketone	ppb	ND	50
35	Methyl methacrylate	ppb	ND	250
37	Methacrylonitrile	ppb	ND	1
38	Methylene chloride	ppb	ND	250
230	2-Nitropropane	ppb	NA	_
39	Pyridine	ppb	ND	25
40.	1,1,1,2-Tetrachloroethane	ppb	ND	50
41	1,1,2,2-Tetrachloroethane	ppb	ND	50
42	Tetrachloroethene	ppb	ND	50
43	Toluene	ppb	D	10
44	Tribromomethane	ppb	ND	50
45	1,1,1-Trichloroethane	ppb	ND	50
46	1,1,2-Trichloroethane	ppb	ND	50
47	Trichloroethene	ppb	ND	50
48	Trichloromonofluoromethane	ppb	ND	50
49	1,2,3-Trichloropropane	ppb	ND	250
231.	1,1,2-Trichloro-1,2,2-trifluoro-	ppb	NA	_
	ethane			
50	Vinyl chloride	ppb	ND	50
215	1,2-Xylene	ppb	D*	50
216	1,3-Xylene	ppb	D*	50
217.	1,4-Xylene	ppb	D*	50
	<u>Semivolatiles</u>			
51	Acenaphthalene	ppm	D	4.600
52.	Acenaphthene	ppm	D	2,500
53	Acetophenone	ррт	ND	3,700
54	2-Acetylaminofluorene	ppm	ND	8,500
55	4-Aminobiphenyl	ppm	ND	5,000
56.	Aniline	ppm	ND	13,000
57.	Anthracene	ppm	D	2,500
58	Aramite	ррп	ND	
59.	Benz(a)anthracene	ppm	D	2,500
218	Benzal chloride	ppm	NA	
60	Benzenethiol	ppm	ND	-
61 <i>.</i>	Deleted	FF'''		
62	Benzo(a)pyrene	ppm	• D	250

⁻ No detection limit available

[^] Analyzed as total xylenes

Table 5-1 (continued)

BDAT			D = Detected	
reference			ND = Not detected	Detection
no	Parameter	Units	NA = Not analyzed	limit
	<u>Semivolatiles</u> (continued)			
63 `	Benzo(b)fluoranthene	ppb	D	250
64	Benzo(ghi)perylene	dqq	ND	5,500
65.	Benzo(k)fluoranthene	ppb	D	250
66.	p-Benzoqu i none	ppb	ND	5,000
67	Bis(2-chloroethoxy)methane	ppb	ND	7,000
6 ð	Bis(2-chloroethyl)ether	ppb	ND	7,500
69	Bis(2-chloroisopropyl)ether	ppb	ND	7,500
70.	Bis(2-ethylhexyl)phthalate	ppb	ND	3,300
71.	4-Bromophenyl phenyl ether	ppb	ND	2,500
72	Butyl benzyl phthalate	ppb	ND	3,300
73	2-sec-Butyl-4,6-dinitrophenol	ppb	NA	-
74.	p-Chloroaniline	ppb	ND	13,000
75	Chlorobenz: late	ppb	ND	5,500
76.	p-Chloro-m-cresol	ppb	ND	5,000
77	2-Chloronaphthalene	ppb	ND	2,500
78.	2-Chlorophenol	ppb	ND	2,500
79	3-Chloropropionitrile	ppb	NA	-
80	Chrysene	ppb	D	2,500
81	ortho-Cresol	ppb	ND	13,000
82	para-Cresol	ppb	ND	13,000
232	Cyclohexanone	ppb	NA	-
83	Dibenz(a,h)anthracene	ppb	ND	3,300
84.	Dibenzo(a,e)pyrene	ppb	ND	2,500
85	Dibenzo(a,i)pyrene	ppb	ND	2,500
86	m-Dichlorobenzene	dad	ИD	2,500
87	o-Dichlorobenzene	ppb	ND	2,500
88	p-Dichlorobenzene	ppb	ND	6,000
89	3,3'-Dichlorobenzidine	ppb	ND	5,000
90	2,4-Dichlorophenol	ppb	ND	3,550
91	2,6-Dichlorophenol	ppb	ND	5,000
92	Diethyl phthalate	ppb	ND	2.50
93.	3,3'-Dimethoxybenzidine	ppb	ND	22,000
94	p-Dimethylaminoazobenzene	ppb	ND	5,00
95	3,3'-Dimethylbenzidine	ppb	ND	5,00
96.	2.4-Dimethylphenol	ppb	ND	3,50
97	Dimethyl phthalate	ppb	ND	2,50
98	Di-n-butyl phthalate	ppb	ND	33,00
99	1,4-Dinitrobenzene	ppb	ND	5,00
100.	4,6-Dinitro-o-cresol	ppb	ND	31,50
100.	2,4-Dinitrophenol	ppb	ND	55,00

Table 5-1 (continued)

BDAT			D = Detected	
reference			ND = Not detected	Detection
no	Parameter	Units	NA = Not analyzed	limit
	<u>Semivolatiles</u> (continued)			
102.	2,4-Dinitrotoluene	ppb	ND	7,500
103	2,6-Dinitrotoluene	dqq	ND	5,00
104	Di-n-octyl phthalate	ppb	ND	3,30
105	Di-n-propylnitrosamine	ppb	ND	2,65
106	Diphenylamine	ppb	ND	5,000
219	Diphenylnitrosamine	ppb	ND	5,00
107	1,2-Diphenylhydrazine	ppb	ND	2,50
108	Fluoranthene	ppb	D	2,50
109.	Fluorene	ppb	D	2,50
110	Hexachlorobenzene	ppb	ND	5,000
111	Hexachlorobutadiene	ppb	ND	8,000
112	Hexachlorocyclopentadiene	ppb	ND	5,000
113	Hexachloroethane	ppb	ND	5,00
114	Hexach lorophene	ppb	NA	,
115	Hexachloropropene	ppb	ND	7,50
116	Indeno(1,2,3-cd)pyrene	ppb	ND	4,90
117	Isosafrole	ppb	ND	5,00
118	Methapyrilene	ppb	ND	16,00
119	3-Methylcholanthrene	dad	ND	4,60
120	4.4'-Methylenebis			,
	(2-chloroaniline)	dqq	ND	5,00
36	Methyl methanesulfonate	dqq	NA	
121	Naphtha lene	dqq	D	2,50
122	1,4-Naphthoguinone	ppb	ND	5,00
123	1-Naphthylamine	dqq	ND	7,50
124	2-Naphthylamine	ppb	ND	17,00
125	p-Nitroaniline	ppb	ND	65,00
126	Nitrobenzene	ppb	ND	2,50
127	4-Nitrophenol	dad	ND	5,00
128.	N-Nitrosodi-n-butylamine	dqq	ND	5,00
129.	N-Nitrosodiethylamine	ppb	ND -	5,00
130.	N-Nitrosodimethylamine	ppb	ND	13,00
131.	N-Nitrosomethylethylamine	dqq	ND	5,00
132.	N-Nitrosomorpholine	ppb	ND	5,00
133.	N-Nitrosopiperidine	ppb	ND	5,00
134	n-Nitrosopyrrolidine	ppb	ND	5,00
135.	5-Nitro-o-toluidine	ppb	ND	5,00
136.	Pentachlorobenzene	ppb	ND	5,00
137.	Pentachloroethane	ppb	ND	5,00
138.	Pentachloronitrobenzene	ppb	ND	5,00

Table 5-1 (continued)

BDAT			D = Detected	
reference			ND = Not detected	Detection
10.	Parameter	Units	NA = Not analyzed	limit
	<u>Semivolatiles</u> (continued)			
139	Pentachlorophenol			
140.	Phenacetin	ppb	D	25
141.	Phenanthrene	ppb	ND	5,000
142.	Pheno 1	ppb	D	2,500
220.	Phthalic anhydride	ppb	D	2,000
143	2-Picoline	ppb	NA	-
144.	Pronamide	ppb	ND	5,000
145.	Pyrene	ppb	ND	5,000
146.	Resorcinol	ppb	D	2,500
147.	Safrole	ppb	ND	5,000
148	1,2,4,5-Tetrachlorobenzene	ppb	ND	5,000
149	2,3,4,6-Tetrachlorophenol	ppb	ND	2,500
150	1,2,4-Trichlorobenzene	ppb	ND	9,000
151.	2,4,5-Trichlorophenol	ppb	ND	2,500
152.	2,4,6-Trichlorophenol	ppb	ND	13,000
153	Tris(2,3-dibromopropyl)	ppb	ND	3,650
	phosphate	•		
		ppb	NA	-
	<u>Metals</u>			
154.	Ant imony			
155.	Arsenic	m	ND	30
156	Barıum	m	D	0 0
157	Beryllium	m	D	0.0
158.	Cadmıum	m	ND	0 5
159.	Chromium (total)	m	D	0.0
221.	Chromium (hexavalent)	m	D	0.0
160	Copper	т	NA	-
161	Lead	m	D	0.0
162.	Mercury	m	D	0.0
163.	Nickel	m	D	0.0
164	Selenium	m	D	7.5
165.	Silver	m	D	2.0
166	Thallium	m	ND	4.5
167.	Vanadıum	m	D	1.5
168.	Zinc	m	D	4.0
		m	D	0.0
	<u>Inorganics</u>			
169.	Cyanide			
170.	Fluoride	ppm	ND	0.2
171.	Sulfide	ppm	D	0.9
		ppm	D	0 5

Table 5-1 (continued)

BDAT			D = Detected	
reference			ND = Not detected	Detection
no	Parameter	Units	NA = Not analyzed	limit
	Organochlorine pesticides			
172.	Aldrin	ppm	ND	7.
173.	a lpha-BHC	ppm	ND	4.
174	beta-BHC	ppm	ND	7
175.	delta-BHC	ppm	ND	7.
176.	gamma-BHC	ppm	ND	5.
177	Chlordane	ppm	ND	100
178	DDD	ppm	ND	15
179.	DDE	ppm	ND	7
180.	DDT	ppm	ND	15
181	Dieldrin	ppm	ND	7.
182.	Endosulfan I	ppm	ND	7.
183.	Endosulfan II	ppm	ND	7.
184.	Endrin	ppm	ND	7
185	Endrin aldehyde	ppm	ND	15
186	Heptachlor	ppm	ND	5.
187.	Heptachlor epoxide	ppm	ND	7.
188	Isodrin	ppm	ND	7.
189.	Kepone	ppm	ND	40
190.	Methoxyclor	ppm	ND	25
191.	Toxaphene	ppm	ND	100
	Phenoxyacetic acid herbicides			
192	2,4-Dichlorophenoxyacetic acid	ppb	ND	0.
193	Silvex	ppb	ND	0.
194.	2,4,5-T	ppb	ND	0.
	Organophosphorous insecticides			
195.	Disulfoton	ppm	ND	0.
196.	Famphur	ppm	NÐ	0.
197.	Methyl parathion	ppm	ND	0
198.	Parathion	ppm	ND	0.
199.	Phorate	ppm	ND	0.
	PCBs			
200	Aroclar 1016	ppm	ND	1,000
201	Aroclor 1221	ppm	ND	1,000
202.	Aroclor 1232	ppm	ND	1,000

Table 5-1 (continued)

BDAT reference			D = Detected ND = Not detected	Detection
no.	Parameter	Units	NA = Not analyzed	limit
	PCBs (continued)			
203.	Aroclor 1242	ppm	ND	1,000
204	Aroclor 1248	ppm	ND	1,000
205.	Aroclor 1254	ppm	ND	300
206	Aroclor 1260	ppm	ND	400
	Dioxins and furans			
207	Hexachlorodibenzo-p-dioxins	ppt	ND	129
208.	Hexachlorodibenzofurans	ppt	ND	87
209.	Pentachlorodibenzo-p-dioxins	ppt	ND	48
210	Pentachlorodibenzofurans	ppt	ND	41
211.	Tetrachlorodibenzo-p-dioxins	ppt	ND	52
212.	Tetrachlorodibenzofurans	ppt	ND	39
213.	2,3,7,8-Tetrachlorodibenzo- p-dioxin	ppt	ND*	-

^{*} Dioxin analyses were not isomer specific, therefore 2,3,7,8-TCDD was analyzed with all tetrachlorodibenzo-p-dioxins.

they were present at or below their detection limits, or that some constituents may be present such that masking or interference has resulted in the inability to detect them. The Agency analyzed for dioxins and furans in the KOOl waste and treatment residuals and did not detect them in any of the waste streams. The Agency has recently become aware of waste characterization data showing that dioxins and furans may be present in some wood preserving wastes. EPA has not had an ample opportunity to evaluate these data. When EPA completes its analysis of available data it will consider the regulation of these constituents. Therefore, the Agency is reserving the standards for dioxins and furans for a later date. Table 5-2 presents the BDAT list constituents that were present in the untreated KOO1-Creosote waste and the ranges of concentrations. Table 5-3 presents similar data for KOO1-PCP. In general, KOOl waste primarily consists of BDAT list semivolatile constituents, with BDAT list metals and volatiles also being present. All BDAT list constituents that were detected in the untreated KOOl waste were considered for regulation unless the constituent was not present at treatable levels or treatment performance data demonstrating effective treatment by BDAT were not available for that constituent in the waste or for a waste judged to be similar.

5.2 BDAT List Constituents Detected in the Treated Waste

The treatment performance data demonstrate that all of the BDAT list organic constituents are significantly reduced by rotary kiln incineration. Specifically, all BDAT list volatile and semivolatile

Table 5-2 Untreated KOO1-Credsote - BDAT List Constituents Detected

BDAT List Constituent	Range of Concentrations (ppm)
Volatiles	
Benzene	51- 83
To luene	100-170
Ethyl Benzene	55- 87
Xylenes	120-170
S em ivolatiles	
Acenaphtha lene	1,000-<4,600
Acenapht hene	15,000-21,000
Anthracene	7,300-15,0 00
Chrysene	4,100- 4,800
luorene	12,000-18,000
Naphthalene	29,000-43,000
Phenanthrene	29,000-41,000
Phenol	2,400- 3,900
yrene	12,000-17,000
<u>Metals</u>	
Arsenic	0.7 - 2 6
Barıum	12 - 150
Cadmium	0 79- 3 5
Chromaum	1.6 - 8.6
Copper	12 - 39
Mercury	0.35- 1 64
Nickel	1 8 - 7 5
ead	37 - 190
Selenium	0.3 - 1 5
[hallium	2.2 - 8 0
/anadium	0.82- 4.0
Zinc	40 - 200

Table 5-3 Untreated K001-PCP - BDAT List Constituents Detected

BDAT List Constituent	Range of Concentrations (ppm)
Volatiles	
To luene	10-39
Semivolatiles	
Acenaphthene	13,000-18,000
Anthracene	8,500-13,000
Benz(a)anthracene	<2,500- 3,400
Benzo(a)pyrene	<250- 940
Benzo(b and/or k)fluoranthrene	940- 2,300
Chrysen e	<2,500- 3,600
Fluoranthene	13,000-21,000
Fluorene	8,200-12,000
Naphtha lene	26,000-43,000
Pentachlorophenol	920- 3,000
Phenanthrene	28,000-42,000
Pyrene	9,200-15,000
<u>Metals</u>	
Arsenic	1 1 - 2 9
Barium	17 - 30
Cadmium	0.4 - 0 6
Chromium	1 5 - 2 7
Copper	6 7 - 11
Lead	6.3 - 11
Mercury	0.0064 - 0 11
Zinc	30 - 64

constituents detected in the untreated waste are reduced to concentrations below their detection limits. Because all of the BDAT list volatiles and semivolatile constituents detected in the untreated waste were reduced to concentrations below their detection limits by rotary kiln incineration, these compounds were regarded as potential regulated constituents, as they are indicators of effective treatment for KOO1 waste.

As explained in Section 1, the Agency is not proposing to regulate all of the BDAT list constituents considered for regulation. In general, the Agency has considered whether some constituents are adequately controlled by the regulation of another constituent. For organic constituents, determination of adequate control was based on an evaluation of the characteristics of the constituents that would affect treatment performance of rotary kiln incineration. Specifically, the waste characteristics affecting performance, as discussed in the incineration discussion in Section 3.2.1, include the volatility (boiling point) and bond dissociation energies of the constituents of a waste. Consistent with the theory of combustion, constituents having higher boiling points and higher bond dissociation energies are the most difficult to destroy. Also, BDAT list organic constituents present in the untreated waste in the highest concentrations are believed to be among the constituents that are most difficult to treat to nondetectable levels.

For KOOl organic treatment by rotary kiln incineration, the constituents proposed for regulation are naphthalene, pentachlorophenol, phenathrene, pyrene, toluene and xylenes (total). These constituents were selected because they are indications of effective treatment and it is believed that other BDAT list organics will be treated to levels equivalent to or lower than these constituents. In general, these constituents are present in the untreated waste in the highest concentrations, have high bond dissociation energies, and/or high boiling points. Table C-1 in Appendix C contains a ranking of the BDAT list organic constituents based on concentration, boiling point, and bond dissociation energy. Specifically, these constituents were selected for the following reasons:

- (1) Napthalene BDAT list constituent present in the untreated waste in the highest concentration.
- (2) Pentachlorophenol Highly chlorinated constituent which will serve as indicator for all chlorinated BDAT list organics.
- (3) Phenanthene BDAT list constituent present in the untreated waste in the highest concentration second to naphthalene, with moderate boiling point, and bond dissociation energy.
- (4) Pyrene BDAT list semivolatile with bond dissociation energy among the highest of any BDAT list semivolatile organic, high boiling point, and present in higher concentrations than other constituents with comparable bond dissociation energies and boiling points.
- (5) Toluene BDAT list volatile organic constituent present in highest concentration.

(6) Xylene - BDAT list volatile constituent present in untreated waste with highest bond dissociation energy and boiling point and concentrations similar to toluene.

EPA believes that the other BDAT list organic constituents present in K001 wastes will be adequately controlled by rotary kiln incineration if these regulated constituents are controlled to concentrations below their detection limits.

Several BDAT metal constituents were detected in the untreated waste collected by the Agency with zinc, lead, barium, and copper being present in the highest concentrations. Rotary kiln incineration is not designed to treat metals, however, metal constituents present in the untreated waste will be present in the incinerator ash, scrubber water and scrubber water treatment nonwastewater residuals. Generally, the metals present in the highest concentrations in the untreated waste are also present in these residuals at the highest concentrations. Whether a BDAT list metal constituent present in the untreated waste will be detected in the ash or the scrubber waste will depend on the volatility of the constituent and the operating temperature of the rotary kiln incinerator. In this case of KOOl as tested by the Agency, the BDAT list metal constituents present in the highest concentration in the untreated waste were also among the constituents most prevalent in both wastewaters and nonwastewater residuals. These constituents, including zinc, lead, barium, and copper, were regarded as potential regulated constituents.

All stabilization data from wastes similar to KOO1 nonwastewaters were examined, and none of the data showed effective treatment for barium. As a result, the three BDAT list metals present in the waste in the highest concentrations, zinc, lead, and copper, were selected as regulated constituents because available stabilization performance showed effective treatment for these constituents. The Agency believes that other BDAT list metals will be adequately controlled by the regulation of these constituents because the others are typically present at significantly lower concentrations.

For the KOO1 scrubber waters, zinc and lead were generally present in the highest concentrations. Several other BDAT list metals including barium, copper, and thallium were also present. EPA examined available treatment performance data for chemical precipitation and filtration for which design and operating data were available. The performance data identified from the Agency's testing at Envirite did not indicate treatment of thallium or barium in wastewaters. However, these data for chemical precipitation and filtration did show effective treatment for lead, copper, and zinc. The Agency believes that the regulation of zinc, lead, and copper, will adequately control other BDAT list metals present in wastewater forms of KOO1.

5.3 <u>Selection of Regulated Constituents</u>

The regulated constituents proposed for KOOl are as follows:

K001 - Nonwastewater

<u> KOO1 - Wastewater</u>

Naphthalene Pentachlorophenol Phenanthrene Pyrene Toluene

Xylenes (total)

Xylenes Copper Lead Zinc Naphthalene Pentachlorophenol Phenanthrene Pyrene Toluene Xylenes (total)

Copper Lead Zinc

6. CALCULATION OF BDAT TREATMENT STANDARDS

The purpose of this section is to present the actual treatment standards for the regulated constituents selected in Section 5. The standards were calculated based on the performance data from the treatment technologies determined in Section 4 to represent BDAT. Included in this section is a detailed discussion of the calculation of treatment standards for the nonwastewater and wastewater forms of KOO1.

As discussed in Section 1, the Agency calculated the BDAT treatment standards for KOO1 by following a four-step procedure: (1) editing the data, (2) correcting the data using recovery data, (3) calculating variability factors, and (4) calculating the actual treatment standards by multiplying the average accuracy corrected composition data by the appropriate variability factor. The four steps in this procedure are discussed in detail in Sections 6.1 through 6.4.

6.1 Editing the Data

6.1.1 BDAT List Organics Treatment

As discussed in Section 3, the Agency collected nine data sets for rotary kiln incineration of KOOl waste at two separate test facilities. The Agency evaluated the nine data sets and determined that the treatment systems were well operated during the sampling periods. These data sets also included the appropriate analytical tests to evaluate treatment performance of incineration. Because incineration is a destruction technology for organics, total constituent concentration is the best measure of performance. The quality assurance/quality control

data were also available for the BDAT list organics and metals, as previously discussed in detail, in Section 4.

6.1.2 BDAT List Metals Treatment

Incineration of KOO1 results in the generation of two treatment residuals: ash (nonwastewater KOO1 residual) and scrubber water (wastewater KOO1 residual). Because the untreated KOO1 waste contains BDAT list metal constituents, these treatment residuals also contain metals at treatable concentrations. As discussed in Section 3, the Agency does not have treatment performance data specifically for BDAT list metals in the wastewater and nonwastewater forms of KOO1. However, EPA does have treatment performance data for wastes that the Agency believes are similar to these KOO1 residuals.

For the wastewater form of KOO1 (scrubber water), treatment performance data were available for the treatment system consisting of chemical precipitation and filtration. Eleven data sets were available for treatment of zinc, lead, and copper containing wastewaters. The design operating data collected for this treatment system indicate that it was well designed and well operated during the time of sampling. In addition, the analytical testing data for total composition of BOAT list metals were the appropriate tests for this technology. However, recovery values are not available for metal spikes and metal spike duplicates from the treatment data transferred from the Onsite Engineering Report for Envirite Co. The recovery data are being transferred from the Onsite Engineering Report for Horsehead Resource Development Co. for KO61. This

is being done because zinc, lead, and copper are present in both wastes and the TCLP extracts from the treated KO61 residual is a similar waste matrix to the wastewaters tested at Envirite.

For nonwastewater residuals containing BDAT list metals requiring stabilization such as nonwastewater residuals from scrubber water treatment by chemical precipitation and filtration or incinerator ash, the Agency identified performance data for zinc, lead, and copper. The treatment data transferred from F006 stabilization data for K001 wastewater treatment nonwastewater residuals and incinerator ash included design and operating data, the appropriate analytical testing to evaluate the performance of stabilization (total composition and TCLP for untreated waste and TCLP for the treated waste), and the required QA/QC analyses. As a result, all of these data were used to develop treatment standards for the BDAT list metals zinc, lead, and copper for nonwastewater forms of K001.

6.2 <u>Correction of Analytical Data</u>

The analytical data used to determine BDAT and calculate treatment standards were adjusted for accuracy in order to take into account the analytical interferences associated with the chemical composition of the sample. This was accomplished by calculating a correction factor from percent recovery data for each regulated constituent. The accuracy adjusted concentration was calculated by multiplying the uncorrected data value by the correction factor. The calculation of corrected data based on recoveries is detailed below for the regulated constituents.

6.2.1 Correction of BDAT List Organics Data

As previously discussed, the BDAT list organic constituents proposed for regulation include naphthalene, pentachlorophenol, phenanthrene, pyrene, toluene, and xylenes (total). All of these constituents were detected in the untreated KOOl waste and were destroyed by rotary kiln incineration to concentrations below their detection limits in the nine data sets collected by the Agency. However, the detection limits attainable for these BDAT list organic constituents in the treatment residuals varied. Generally, where this occurred, the Agency selected the highest detection limit measured for each regulated constituent in each waste matrix (wastewater and nonwastewater). The treatment standards were developed using these high detection limit because lower detection limits may not be consistently achievable. The treatment performance data for the proposed regulated organic presented in this section and used to calculate the treatment standards reflect this change.

The recovery data used to develop accuracy-corrected data for BDAT list organics in K001 treatment residuals were from the accompanying matrix spike recoveries from the data with the higher detection limits. As noted in Section 4, it was necessary in one instance to transfer recovery data for pentachlorophenol in the scrubber water. The detection limit used to calculate the standard was from the treatment of the K001-pentachlorophenol waste, while the recovery data used were transferred from the matrix spike recovery performed for pentachlorophenol in the scrubber water from the K001-creosote waste.

The accuracy correction factors used for all regulated organic constituents in both the wastewater and nonwastewater residuals are summarized in Table 6-1. The matrix spike recovery data used to correct the data used in calculating the treatment standards are presented in Appendix B.

6.2.2 Correction of BDAT List Metals Data

For KOOl nonwastewater residuals the Agency is proposing treatment standards for zinc, lead, and copper. As stated previously, the performance data for stabilization of these constituents were transferred from FOO6 stabilization data. As a result, the matrix spike recovery values used to correct the data were developed with the analytical data for those FOO6 wastes. The correction factors used and calculations of the corrected values for the proposed regulated BDAT list metals in the nonwastewater residuals are presented in Table 6-2.

The Agency is proposing to regulate zinc, lead, and copper in the KOO1 wastewater residual (scrubber water) from rotary kiln incineration. The treatment performance data for treatment by chemical precipitation and filtration were transferred from the Onsite Engineering Report for Envirite. The recovery data were transferred from the Onsite Engineering Report for Horsehead Resource Development Co. for KO61. The correction of the analytical data and the correction factors used are presented in Table 6-3.

Table 6-1 Corrected Values for BDAT List Organics

BDAT List Constituent	Uncorrected Detection Limit (ppm)	Correction Factor	Accuracy Corrected Value	
Naphthalene (ash)	<2.5	1.14	2.85	
Naphthalene (water)	< 0 050	1.06	0.053	
Pentachlorophenol (ash)	<12 5	1 05	13.125	
Pentachlorophenol (water)	<0.250	1.25	0.313	
Phenanthrene (ash)	<2.5	1.14	2.85	
Phenanthrene (water)	<0.050	1.06	0 053	
Pyrene (ash)	<2.5	1.04	2.60	
Pyrene (water)	<0.050	1 00	0.050	
Toluene (ash)	<0.050	1.01	0.051	
Toluene (water)	<0.050	1.01	0.051	
Xylenes (ash)	<0.050	1.16	0.058	
Xylenes (water)	< 0 050	1.15	0.058	

Table 6-2 Corrected Values for Regulated Metal Constituents
Treated by Stabilization

Constituent		Accurac				Mean (ppm)				
	1	2	3	4	5	6	7	8	9	.,,,
BDAT List <u>Metals</u>										
opper	-	0 27	-	0.16	0 29	0.31	0 45	0 35	0.50	0.33
.ead	-	0 39	0 34	0 23	0 37	0.39	0 41	0 40	0.29	0.35
inc	0.03	0 01	0.05	0 01	0 04	0.03	0 02	0.02	0.01	0.024

Table 6-3 Corrected Values for Regulated Metal Constituents Treated by Chemical Precipitation and Filtration

	Accuracy-corrected concentration (mg/l) CorrectionSample Set #												
Constituent	factor	1	2	3	4	5	6	7	8	9	10	11	Mean (mg/1)
BDAT Metals						,							
Copper	1 20	0.25	0.18	0.25	0.08	0.17	0 14	0.19	0.19	0.10	0.17	0 29	0.18
Lead	1.32	<0.013	<0.013	<0.013	<0.013	<0.013	<0.013	<0.013	<0.013	<0.013	<0.013	<0.013	0.013
Zinc	1.02	0 128	0.117	0.143	1.653	0 128	0 097	0.117	0.133	0.061	0.071	0 102	0.25

6.3 <u>Calculation of Variability Factors</u>

The variability factor represents the variability inherent in the treatment process and the sampling and analytical methods. Variability factors are calculated based on the treatment data for each of the regulated constituents. The general methodology for calculating variablity factors is presented in Appendix A. In cases where all of the treated values for a constituent are below the detection limits, a detection limit of 2.8 was used. The methodology used to calculate this variability factor is presented in Appendix A. The variability factors calculated for the regulated constituents for KOO1 are presented in Table 6-4. Appendix E of this report presents these calculations.

6.4 <u>Calculation of Treatment Standards</u>

The treatment standards for the proposed regulated constituents were calculated by multiplying the average accuracy corrected values by the appropriate variability factor. The proposed treatment standards for BDAT are presented in Table 6-4.

Table 6-4 Calculation of Treatment Standards for K001

Regulated Constituent	Average Treated Concentration (mg/kg, mg/l)	Variability Factor (VF)	Treatment Standard (VF x Average)
Naphthalene (ash)	2.85	2.8	7 98
Naphthalene (water)	0 053	2 8	0 148
Pentachlorophenol (ash)	13.13	2.8	36 75
Pentachlorophenol (water)	0.313	2 8	0 875
Phenanthrene (ash)	2 85	2.8	7.98
Phenanthrene (water)	0 053	2 8	0 148
Pyrene (ash)	2 60	2.8	7.28
Pyrene (water)	0.050	2 8	0.140
Toluene (ash)	0 051	2.8	0 143
Toluene (water)	0 051	2 8	0 143
Xylenes (ash)	0.058	2.8	0.16
Xyle nes (w ater)	0 058	2 8	0.16
Copper (nonwastewater)	0.33	2 2	0 71
Copper (wastewater)	0.18	2.31	0.42
Lead (nonwastewater)	0 35	1 5	0 53
Lead (wastewater)	0.013	2.8	0 037
Zinc (nonwastewater)	0.024	3.6	0.086
Zinc (wastewater)	0.250	4 13	1.0

REFERENCES

Ackerman D.G., McGaughey J.F., Wagoner D.E., "At Sea Incineration of PCB-Containing Wastes on Board the M/T Vulcanus," USEPA, 600/7-83-024, April 1983.

Acurex Corporation, 1982. Emissions and Residue Values from Waste Disposal during Wood Preserving (as cited by <u>Summary of Available Waste Composition Data from Review of Literature and Data Bases for Use in Treatment Technology Application and Evaluation for "California List" <u>Waste Streams</u>. Prepared for EPA Office of Solid Waste by Versar, Inc. under Contract No. 68-01-7053, Work Assignment No. 38, April 1986).</u>

Ajax Floor Products Corp. n.d. Product literature: technical data sheets, Hazardous Waste Disposal System. P.O. Box 161, Great Meadows, N.J. 07838.

Aldrich, James R. 1985. "Effects of pH and proportioning of ferrous and sulfide reduction chemicals on electroplating waste treatment sludge production." In <u>Proceeding of the 39th Purdue Industrial Waste Conference</u>, May 8, 9, 10, 1984. Stoneham, MA: Butterworth Publishers.

Austin, G.T. 1984. Shreve's chemical process industries, 5th ed. New York: McGraw-Hill

Bishop, P.L., Ransom, S.B., and Grass, D.L. 1983. Fixation Mechanisms in Solidification/Stabilization of Inorganic Hazardous Wastes. In Proceedings of the 38th Industrial Waste Conference, 10-12 May 1983, at Purdue University, West Lafayette, Indiana.

Bonner TA, et al., Engineering Handbook for Hazardous Waste Incineration. SW-889. Prepared by Monsanto Research Corporation for U.S. EPA NTIS PB 81-248163. June 1981.

Brown, K.W. and Associates, 1981. Hazardous Waste Land Treatment (as cited by Summary of Available Waste Composition Data from Review of Literature and Data Bases for Use in Treatment Technology Application and Evaluation for "California List" Waste Streams. Prepared for EPA Office of Solid Waste by Versar, Inc. under Contract No. 68-01-7053, Work Assignment No. 38, April 1986).

Castaldini C., et al., <u>Disposal of Hazardous Wastes in Industrial Boilers on Furnaces</u>, Noyes Publications, New Jersey, 1986.

Chemical Waste Management. 1987. Technical Report 87-117.

REFERENCES - (continued)

Cherry, Kenneth F. 1982. <u>Plating Waste Treatment</u>. Ann Arbor, MI; Ann Arbor Science, Inc. pp 45-67.

Cushnie, George C., Jr. 1984. <u>Removal of Metals from Wastewater:</u>
<u>Neutralization and Precipitation</u>. Park Ridge, NJ; Noyes Publications.
pp 55-97.

Conner, J.R. 1986. Fixation and Solidification of Wastes. Chemical Engineering. Nov. 10, 1986.

Cullinane, M.J., Jr., Jones, L. W.., and Malone, P.G. 1986. Handbook for stabilization/solidification of hazardous waste. U.S. Army Engineer Waterways Experiment Station. EPA report No. 540/2-86/001. Cincinnati, Ohio: U.S. Environmental Protection Agency.

Cushnie, George C., Jr. 1985. <u>Electroplating Wastewater Pollution</u> <u>Control Technology</u>. Park Ridge, NJ; Noyes Publications. pp 48-62, 84-90.

DPRA, 1984. Data from OSW Mail Survey: Generator Questionnaire, Incinerator Questionnaire, Land Disposal Questionnaire. (as cited by Summary of Available Waste Composition Data from Review of Literature and Data Bases for Use in Treatment Technology Application and Evaluation for "California List" Waste Streams. Prepared for EPA Office of Solid Waste by Versar, Inc. under Contract No. 68-01-7053, Work Assignment No. 38, April 1986).

Eckenfelder, W.W. 1985. Wastewater treatment. <u>Chemical Engineering</u>, 85:72.

Electric Power Research Institute. 1980. FGD sludge disposal manual, 2nd ed. Prepared by Michael Baker, Jr., Inc. EPRI CS-1515 Project 1685-1 Palo Alto, California: Electric Power Research Institute.

Federal Register. 1986. Hazardous Waste Management systems; Land Disposal Restrictions; Final Rule; Appendix I to Part 268 - Toxicity Leaching Procedure (TCLP). Vol. 51, No. 216. November 7, 1986 pp. 40643-40654.

Grain, Richard W. 1981. Solids removal and concentration. In <u>Third Conference on Advanced Pollution Control for the Metal Finishing Industry</u>. Cincinnati, Ohio: U.S. Environmental protection Agency. pp. 56-62.

Gurnham, C.F. 1955. <u>Principles of Industrial Waste Treatment</u>. New York; John Wiley and Sons. pp 224-234.

REFERENCES - (continued)

Handbook of Industrial Waste Compositions in California, 1978. (as cited by <u>Summary of Available Waste Composition Data from Review of Literature and Data Bases for Use in Treatment Technology Application and Evaluation for "California List" Waste Streams. Prepared for EPA Office of Solid Waste by Versar, Inc. under Contract No. 68-01-7053, Work Assignment No. 38, April 1986).</u>

Illinois EPA, 1983. Special Waste Authorization File. (as cited by Summary of Available Waste Composition Data from Review of Literature and Data Bases for Use in Treatment Technology Application and Evaluation for "California List" Waste Streams. Prepared for EPA Office of Solid Waste by Versar, Inc. under Contract No. 68-01-7053, Work Assignment No. 38, April 1986).

Kirk-Othmer. 1980. Encyclopedia of Chemical Technology, 3rd ed., "Flocculation", Vol. 10. New York; John Wiley and Sons. pp 489-516.

Lanouette, Kenneth H. 1977. "Heavy metals removal." <u>Chemical</u> <u>Engineering</u>, October 17, 1977, pp. 73-80.

Mishuck, E. Taylor, D.R., Telles, R. and Lubowitz, H. 1984. Encapsulation/Fixation (E/F) mechanisms. Report No. DRXTH-TE-CR-84298. Prepared by S-Cubed under Contract No. DAAK11-81-C-0164..

Mitre Corporation, 1981. Composition of Hazardous Waste Streams. (as cited by <u>Summary of Available Waste Composition Data from Review of Literature and Data Bases for Use in Treatment Technology Application and Evaluation for "California List" Waste Streams. Prepared for EPA Office of Solid Waste by Versar, Inc. under Contract No. 68-01-7053, Work Assignment No. 38, April 1986).</u>

Mitre Corp. "Guidance Manual for Hazardous Waste Incinerator Permits." NTIS PB84-100577. July 1983.

Myers, L.H., et al., 1979. <u>Indicatory Fate Study</u>, EPA OGC2-78-179. (as cited by <u>Summary of Available Waste Composition Data from Review of Literature and Data Bases for Use in Treatment Technology Application and Evaluation for "California List" <u>Waste Streams</u>. Prepared for EPA Office of Solid Waste by Versar, Inc. under Contract No. 68-01-7053, Work Assignment No. 38, April 1986).</u>

Novak RG, Troxler WL, Dehnke TH, "Recovering Energy from Hazardous Waste Incineration," Chemical Engineering Progress 91:146 (1984).

- REFERENCES (continued)
- Oppelt ET, "Incineration of Hazardous Waste"; JAPCA; Volume 37, No. 5; May 1987.
- Patterson, James W. 1985. <u>Industrial Wastewater Treatment Technology</u>, 2nd Ed. Butterworth Publishers; Stoneham, MA.
- Perry, Robert H. and Chilton, Cecil H. 1973. <u>Chemical Engineers'</u> Handbook. Fifth Edition. New York: McGraw Hill, Inc., Section 19.
- Pojasek RB. 1979. "Solid-Waste Disposal: Solidification" <u>Chemical Engineering</u> 86(17): 141-145.
- Rudolfs, William. 1953. <u>Industrial Wastes</u>. <u>Their Disposal and Treatment</u>. L.E.C. Publishers Inc., Valley Stream, NY. p. 294
- Santoleri J.J., "Energy Recovery-A By-Product of Hazardous Waste Incineration Systems," in Proceedings of the 15th Mid-Atlantic Industrial Waste Conference on Toxic and hazardous Waste, 1983.
- U.S. Department of Commerce, 1982 Census of Manufacturers Miscellaneous Chemical Products. December 1984.
- USEPA. 1980. U.S. Environmental Protection Agency. RCRA Listing Background Document Waste Code K001.
- USEPA. 1980a. U.S. Environmental Protection Agency. U.S. Army Engineer Waterways Experiment Station. Guide to the disposal of chemically stabilized and solidified waste. Prepared for HWERL/ORD under Interagency Agreement No. EPA-IAG-D4-0569. PB81-181505. Cincinnati, Ohio.
- USEPA. 1983. <u>Treatability Manual</u>, Volume III, Technology for Control/Removal of Pollutants. EPA-600/2-82-001C, January 1983. pp 111.3.1.3-2.
- USEPA. 1985. Characterization of Waste Streams Listed in 40 CFR; Section 261, Waste Profiles. Prepared for the Waste Identification Branch, Characterization and Assessment Division, U.S. EPA. Prepared by Environ Corporation, Washington, D.C. 1985.
- USEPA. 1986. Onsite Engineering Report of Treatment Technology Performance and Operation for Envirite Corporation. York, Pennsylvania. Washington, D.C.: U.S. Environmental Protection Agency.

REFERENCES - (continued)

USEPA. 1986. Test Methods for Evaluating Solid Waste; physical/chemical methods. Third Edition. U.S. EPA. Office of Solid Waste and Emergency Response. November 1986.

USEPA. 1988. Onsite Engineering Report for Horsehead Resource Development Co., Inc. Palmerton, Pennsylvania for K061. Washington, D.C.: U.S. Environmental Protection Agency.

U.S. EPA. 1987. United States Environmental Protection Agency, Office of Solid Waste. Onsite Engineering Report for K001 - Creosote. November 23, 1987.

U.S. EPA. 1987. United States Environmental Protection Agency, Office of Solid Waste. Onsite Engineering Report for K001 - PCP. November 12, 1987.

Versar Inc. 1984. Estimating PMN Incineration Results (Draft). U.S. Environmental Protection AGency, Exposure Evaluation Division, Office of Toxic Substances, Washington, DC. EPA Contract No. 68-01-6271, Task No. 66.

Vogel G, et al., "Incineration and Cement Kiln Capacity for Hazardous Waste Treatment," in Proceedings of the 12th Annual Research Symposium. Incineration and Treatment of Hazardous Wastes. Cincinnati, Ohio. April, 1986.

APPENDIX A

APPENDIX A

A.1 F Value Determination for ANOVA Test

As noted earlier in Section 1.0, EPA is using the statistical method known as analysis of variance in the determination of the level of performance that represents "best" treatment where more than one technology is demonstrated. This method provides a measure of the differences between data sets. If the differences are not statistically significant, the data sets are said to be homogeneous.

If the Agency found that the levels of performance for one or more technologies are not statistically different (i.e., the data sets are homogeneous), EPA would average the long term performance values achieved by each technology and then multiply this value by the largest variability factor associated with any of the acceptable technologies. If EPA found that one technology performs significantly better (i.e., the data sets are not homogeneous), BDAT would be the level of performance achieved by the best technology multiplied by its variability factor.

To determine whether any or all of the treatment performance data sets are homogeneous using the analysis of variance method, it is necessary to compare a calculated "F value" to what is known as a "critical value." (See Table A-1.) These critical values are available in most statistics texts (see, for example, <u>Statistical Concepts and Methods</u> by Bhattacharyya and Johnson, 1977, John Wiley Publications, New York).

Where the F value is less than the critical value, all treatment data sets are homogeneous. If the F value exceeds the critical value, it is necessary to perform a "pair wise F" test to determine if any of the sets are homogeneous. The "pair wise F" test must be done for all of the various combinations of data sets using the same method and equation as the general F test.

The F value is calculated as follows:

- (i) All data are natural logtransformed.
- (ii) The sum of the data points for each data set is computed (T_i) .
- (iii) The statistical parameter known as the sum of the squares between data sets (SSB) is computed:

$$SSB = \begin{bmatrix} k & T_i^2 \\ \sum_{i=1}^{K} \left(\frac{T_i^2}{n_i}\right) \end{bmatrix} - \begin{bmatrix} k & T_i \\ \sum_{i=1}^{K} T_i \end{bmatrix}^2$$

where:

k = number of treatment technologies $n_{\dot{1}}$ = number of data points for technology i N = number of data points for all technologies $T_{\dot{1}}$ = sum of natural logtransformed data points for each technology.

(iv) The sum of the squares within data sets (SSW) is computed:

$$SSW = \begin{bmatrix} k & n_i \\ \sum & \sum \\ i=1 & j=1 \end{bmatrix} \times x^2_{i,j} - \sum_{i=1}^{k} \left(\frac{T_i^2}{n_i} \right)$$

where:

 $x_{i,j}$ = the natural logtransformed observations (j) for treatment technology (i).

- (v) The degrees of freedom corresponding to SSB and SSW are calculated. For SSB, the degree of freedom is given by k-1. For SSW, the degree of freedom is given by N-k.
- (vi) Using the above parameters, the F value is calculated as follows:

$$\cdot MSB$$

$$F = MSW$$

where:

MSB = SSB/(k-1) and MSW = SSW/(N-k).

A computational table summarizing the above parameters is shown below.

Computational Table for the F Value

Source	Degrees of freedom	Sum of squares	Mean square	F
Between	K-1	SSB	MSB = SSB/k-1	MSB/MSW
Within	N-k	SSW	MSW = SSW/N-k	

Below are three examples of the ANOVA calculation. The first two represent treatment by different technologies that achieve statistically similar treatment; the last example represents a case where one technology achieves significantly better treatment than the other technology.

Example 1 Methylene Chloride

	Steam stripping				Biological trea	tmont	
Influent (µg/l)	Effluent (μg/l)	ln(effluent)	[ln(effluent)] ²	Influent (μg/1)	Effluent (µg/l)	ln(effluent)	[ln(effluent)] ²
1550 00	10 00	2 30	5.29	1960.00	10.00	2 30	5 29
1290 00	10 00	2.30	5 29	2568.00	10.00	2.30	5 29 5 29
1640 00	10 00	2.30	5.29	1817.00	10 00	2.30	5 29
5100 00	12.00	2 48	6.15	1640.00	26.00	3.26	10 63
1450.00	10.00	2.30	5.29	3907.00	10.00	2.30	5.29
4600 00	10.00	2 30	5.29	0007.00	10.00	2.30	J. £ 3
1760 00	10.00	2.30	5.29				
2400.00	10.00	2.30	5.29				
4800 00	10.00	2.30	5 29				
12100 00	10.00	2 30	5 29				
Sum -	-	23.18	53.76	_	_	12 46	31 79
Sample Siz	e:						
10	10	10	-	5	5	5	-
Mean							
3669	10.2	2.32	-	2378	13.2	2.49	-
Standard D	eviation						
3326.67	63	.06	-	923.04	7.15	43	-
Variabilit	y Factor.						
	1.15	-	-	-	2.48	-	-

ANOVA Calculations.

$$SSB = \begin{bmatrix} k \\ \sum_{i=1}^{K} \left(\frac{T_{i}^{2}}{n_{i}} \right) \end{bmatrix} \begin{bmatrix} \left[\frac{k}{\sum_{i=1}^{K} T_{i}} \right]^{2} \\ \frac{k}{N} \end{bmatrix}$$

$$SSW = \begin{bmatrix} k & n_{i} \\ \sum_{i=1}^{K} \sum_{j=1}^{N} x^{2}_{i,j} \end{bmatrix} - \sum_{i=1}^{K} \left[\frac{T_{i}^{2}}{n_{i}} \right]$$

MSB = SSB/(k-1)

MSW = SSW/(N-k)

1790g

Example 1 (continued)

F = MSB/MSW

where.

k = number of treatment technologies

 $n_i = number of data points for technology 1$

N = number of natural log transformed data points for all technologies

T = sum of log transformed data points for each technology

 $X_{j,j}$ = the nat log transformed observations (j) for treatment technology (i)

$$n_1 = 10$$
, $n_2 = 5$, $N = 15$, $k = 2$, $T_1 = 23$ 18, $T_2 = 12.46$, $T = 35$ 64, $T^2 = 1270$ 21

$$T_1^2 = 537.31$$
 $T_2^2 = 155.25$

$$SSB = \left(\frac{537.31}{10} + \frac{155.25}{5}\right) - \frac{1270.21}{15} = 0.10$$

SSW =
$$(53.76 + 31.79) - \left(\frac{537.31}{10} + \frac{155.25}{5}\right)$$
 = 0.77

$$MSB = 0.10/1 = 0.10$$

$$MSW = 0.77/13 = 0.06$$

$$F = \frac{0.10}{0.06} = 1.67$$

ANOVA Table

Source	Degrees of	SS	MS	E
	7,0000			'
Between(B)	1	0.10	0 10	1.67
Within(W)	13	0 77	30 0	

The critical value of the F test at the 0.05 significance level is 4.67. Since the F value is less than the critical value, the means are not significantly different (i.e., they are homogeneous).

Note All calculations were rounded to two decimal places Results may differ depending upon the number of decimal places used in each step of the calculations.

Example 2 Trichloroethylene

7	team stripping				<u>Biological trea</u>	itment	
Influent (µg/l)	Effluent (μg/l)	<pre>ln(effluent)</pre>	[ln(effluent)] ²	Influent $(\mu g/1)$	Effluent (μg/l)	ln(effluent)	[ln(effluent)] ²
1650 00	10.00	2.30	5.29	200 00	10 00	2 30	5 29
5200 00	10 00	2 30	5.29	224.00	10.00	2.30	5 29
5000.00	10 00	2 30	5.29	134.00	10 00	2.30	5.29
1720 00	10 00	2.30	5.29	150.00	10.00	2.30	5 29
1560 00	10 00	2 30	5.29	484.00	16.25	2.79	7 78
10300 00	10.00	2.30	5.29	163.00	10.00	2.30	5 29
210.00	10.00	2 30	5.29	182.00	10 00	2.30	5.29
1600.00	27 00	3.30	10.89				
204 00	85 00	4 44	19.71				
160 00	10 00	2 30	5 29				
Sum -	-	26.14	72 92	-	-	16 59	39.52
Sample Size	10	10	-	7	7	7	-
Mean				220	10 89	2 37	-
Me an 2760	19.2	2.61	-	220	10 03	2 0,	
Standard Dev	/lation		-				
2760		2.61	-	120.5	2.36	. 19	-
2760 Standard Dev	viation 23 7		-				-

ANOVA Calculations:

$$SSB = \begin{bmatrix} k \\ \Sigma \\ 1=1 \end{bmatrix} \begin{bmatrix} T_1^2 \\ \overline{n_1} \end{bmatrix} \begin{bmatrix} \begin{bmatrix} k \\ \Sigma \\ 1=1 \end{bmatrix}^2 \end{bmatrix}$$

$$SSW = \begin{bmatrix} k \\ \Sigma \\ 1=1 \end{bmatrix} \begin{bmatrix} n_1 \\ 1=1 \end{bmatrix} \times \begin{bmatrix} N \\ 1=1 \end{bmatrix} \begin{bmatrix} N \\ 1=1 \end{bmatrix} \begin{bmatrix} T_1^2 \\ \overline{n_1} \end{bmatrix}$$

MSB = SSB/(k-1)

MSW = SSW/(N-k)

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Example 2 (continued)

F = MSB/MSW

where.

k = number of treatment technologies

n = number of data points for technology i

N = number of data points for all technologies

T = sum of natural log transformed data points for each technology

 X_{11} = the natural log transformed observations (j) for treatment technology (i)

$$N_1 = 10$$
, $N_2 = 7$, $N = 17$, $k = 2$, $T_1 = 26.14$, $T_2 = 16.59$, $T = 42.73$, $T^2 = 1825.85$, $T_1^2 = 683.30$,

$$T_2^2 = 275.23$$

$$SSB = \begin{cases} 683.30 \\ 10 \end{cases} + \frac{275.23}{7} - \frac{1825.85}{17} = 0.25$$

SSW =
$$(72.92 + 39 52) - \left(\frac{683 30}{10} + \frac{275.23}{7}\right) = 4.79$$

$$MSB = 0.25/1 = 0.25$$

$$MSW = 4.79/15 = 0.32$$

$$F = \frac{0.25}{0.32} = 0.78$$

ANOVA Table

Source	Degrees of freedom	SS	MS	F
Between(B) Within(W)	1 15	0.25 4.79	0.25 0.32	0.78

The critical value of the F test at the 0.05 significance level is 4.54. Since the F value is less than the critical value, the means are not significantly different (i.e., they are homogeneous).

Note: All calculations were rounded to two decimal places. Results may differ depending upon the number of decimal places used in each step of the calculations.

٠. .

Example 3 Chlorobenzene

nfluent (μg/l)	Effluent (μg/l)	<pre>ln(effluent)</pre>	[ln(effluent)] ²	Influent (μg/l)	Effluent (µg/l)	<pre>ln(effluent)</pre>	<pre>ln[(effluent)]²</pre>
7200 00	80.00	4.38	19.18	9206 00	1083.00	6.99	48 8 6
6500 00	70.00	4.25	18.06	16646 00	709.50	6.56	43 03
6075 00	35.00	3.56	12.67	49775.00	460.00	6 13	37 58
3040 00	10.00	2.30	5.29	14731.00	142.00	4.96	24 60
				3159 00	603 00	6 40	40 9€
				€756. 0 0	153 00	5.03	25 30
				3040.00	17.00	2.83	8 01
ium -	_	14 49	55.20	-	-	38.90	228 34
Sample Size:				-	~	7	
4	4	4	-	7	7	7	-
Mean:							
5703	49	. 3 62	-	14759	452.5	5 56	-
Standard Dev	nation:		•				
1835 4	32.24	95		16311.86	379.04	1 42	-
1030 4							
/ariability	Factor						

ANOVA Calculations

$$SSB = \begin{bmatrix} k \\ \sum_{i=1}^{K} \left(\frac{T_i^2}{n_i} \right) \end{bmatrix} \begin{bmatrix} \left[\frac{k}{\sum_{i=1}^{K} T_i} \right]^2 \\ \frac{k}{N} \end{bmatrix}$$

$$SSW = \begin{bmatrix} k & n_1 \\ \sum_{i=1}^{K} \sum_{j=1}^{K} x^2_{i,j} \end{bmatrix} - \frac{k}{N} \begin{bmatrix} \frac{T_i^2}{n_1} \end{bmatrix}$$

MSB = SSB/(k-1)

MSW = SSW/(N-k)

F = MSB/MSW

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Example 3 (continued)

where.

$$k$$
 = number of treatment technologies

$$n_1$$
 = number of data points for technology 1

N = number of data points for all technologies

T = sum of natural log transformed data points for each technology

 $X_{1,j}$ = the natural log transformed observations (j) for treatment technology (i)

$$N_1 = 4$$
, $N_2 = 7$, $N = 11$, $k = 2$, $T_1 = 14$ 49, $T_2 = 38$ 90, $T = 53$ 39, $T^2 = 2850.49$, $T_1^2 = 209.96$

$$T_2^2 = 1513.21$$

$$SSB = \begin{cases} 209.96 & + & \frac{1513.21}{7} \\ & & \end{cases} - \frac{2850 \ 49}{11} = 9.56$$

$$SSW = (55.20 + 228.34) - \left[\frac{209.96}{4} + \frac{1513.21}{7} \right] = 14.88$$

$$MSB = 9.52/1 = 9.52$$

$$MSW = 14.88/9 = 1.65$$

$$F = 9.52/1.65 = 5.77$$

ANOVA Table

Source	Degrees of freedom	SS	MS	F
Jetween(B)	1	9.53	9.53	5.77
lithin(W)	9	14.89	1.65	

The critical value of the F test at the 0.05 significance level is 5.12 Since the F value is larger than the critical value, the means are significantly different (i.e., they are heterogeneous).

Note: All calculations were rounded to two decimal places. Results may differ depending upon the number of decimal places used in each step of the calculations.

A.2. Variability Factor

 $VF = \frac{C_{99}}{Mean}$

where:

VF = estimate of daily maximum variability factor determined from a sample population of daily data.

 C_{99} = Estimate of performance values for which 99 percent of the daily observations will be below. C_{99} is calculated using the following equation: C_{99} = Exp(y + 2.33 Sy) where y and Sy are the mean and standard deviation, respectively, of the logtransformed data.

Mean = average of the individual performance values.

EPA is establishing this figure as an instantaneous maximum because the Agency believes that on a day-to-day basis the waste should meet the applicable treatment standards. In addition, establishing this requirement makes it easier to check compliance on a single day. The 99th percentile is appropriate because it accounts for almost all process variability.

In several cases, <u>all</u> the results from analysis of the residuals from BDAT treatment are found at concentrations less than the detection limit. In such cases, all the actual concentration values are considered unknown and hence, cannot be used to estimate the variability factor of the analytical results. Below is a description of EPA's approach for calculating the variability factor for such cases with all concentrations below the detection limit.

It has been postulated as a general rule that a lognormal distribution adequately describes the variation among concentrations. As a general rule for the BDAT program, empiric observations on

concentrations for several constituents showed that the treatment residual concentrations were distributed approximately lognormally. Therefore, the lognormal model has been used routinely in the EPA development of numerous regulations in the Effluent Guidelines program and in the BDAT program. The variability factor (VF) was defined as the ratio of the 99th percentile (C_{qq}) of the lognormal distribution to its arithmetic mean (Mean).

$$VF = \frac{C_{99}}{Mean} \tag{1}$$

The relationship between the parameters of the lognormal distribution and the parameters of the normal distribution created by taking the natural logarithms of the lognormally-distributed concentrations can be found in most mathematical statistics texts (see for example: Distribution in Statistics-Volume 1 by Johnson and Kotz, 1970). of the lognormal distribution can be expressed in terms of the mean (μ) and standard deviation (σ) of the normal distribution as follows:

$$C_{99} = Exp (\mu + 2.33\sigma)$$
 (2)
Mean = $Exp (\mu + .5\sigma^2)$ (3)

$$Mean = Exp (\mu + .5\sigma^2)$$
 (3)

Substituting (2) and (3) in (1) the variability factor can then be expressed in terms of σ as follows:

$$VF = Exp (2.33 \sigma - .5\sigma^{2})$$
 (4)

For residuals with concentrations that are not all below the detection limit, the 99 percentile and the mean can be estimated from the actual analytical data and accordingly, the variability factor (VF)

can be estimated using equation (1). For residuals with concentrations that are below the detection limit the following steps demonstrated the approach that is used to approximate the value of σ and, hence, calculate the VF using equation (4).

Step 1: The actual concentrations follow a lognormal distribution (truncated). The upper limit (UL) is equal to the detection limit. The lower limit (LL) is assumed to be equal to one tenth of the detection limit. This assumption is based on the fact that data from well-designed and well-operated treatment systems generally falls within one order of magnitude.

Step 2: The natural logarithms of the concentrations have a normal distribution (truncated) with an upper limit equal to ln (UL) and a lower limit equal to ln (LL).

Step 3: The standard deviation (σ) of the normal distribution is approximated by

$$\sigma$$
 = [(ln (UL) - ln (LL)] / [(2)(2.33)] = [ln(UL/LL)] / 4.66
when LL = (0.1)(UL) then σ = (ln10) / 4.66 = 0.494

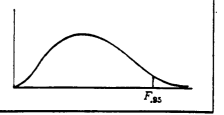
Step 4: Substitution of the value from Step 3 in equation (4) yields the variability factor, VF.

VF = 2.8

Table A-1

95th PERCENTILE VALUES FOR THE \vec{F} DISTRIBUTION

 n_1 = degrees of freedom for numerator n_2 = degrees of freedom for denominator (shaded area = .95)



712	1	2	3	4	5	6	8	12	16	20	30	40	50	100	•
1	161.4	199.5	215.7	224.6	230.2	234.0	238.9	243.9	246.3	248.0	250.1	251.1	252.2	253.0	254.3
2	18.51	19.00	19.16	19.25	19.30	19.33	19.37	19.41	19.43	19.45	19.46	19.46	19.47	19.49	19.50
3	10.13	9.55	9.28	9.12	9.01	8.94	8.85	8.74	8.69	8.66	8.62	8.6 0	8.58	8.56	8.53
4	7.71	6.94	6.59	6.39	6.26	6.16	6.04	5.91	5.84	5.80	5.75	5.71	5.70	5.66	5.63
5	6.61	5.79	5.41	5.19	5.05	4.95	4.82	4.68	4.60	4.56	4.50	4.46	4.44	4.40	4.36
6	5.99	5.14	4.76	4.53	4.39	4.28	4.15	4.00	3.92	3.87	3.81	3.77	3.75	3.71	3.67
7	5.59	4.74	4.35	4.12	3.97	3.87	3.73	3.57	3.49	3.44	3.38	3.34	3.32	3.28	3.23
8	5.32	4.46	4.07	3.84	3.69	3.58	3.44	3.28	3.20	3.15	3.08	3.05	3.03	2.98	2.93
9	5.12	4.26	5.86	3.63	3.48	3.37	3.23	3.07	2.98	2.93	2.86	2.82	2.80	2.76	2.71
10	4.96	4.10	3.71	3.48	3.33	3.22	3.07	2.91	2.82	2.77	2.70	2.67	2.64	2.59	2.54
11	4.84	3.98	3.59	3.36	3.20	3.09	2.95	2.79	2.70	2.65	2.57	2.53	2.50	2.45	2.40
12	4:75	3.89	3.49	3.26	3.11	3.00	2.85	2.69	2.60	2.54	2.46	2.42	2.40	2.35	2.30
13	4.67	3.81	3.41	3.18	3.03	2.92	2.77	2.60	2.51	2.46	2.38	2.34	2.32	2.26	2.21
14	4.60	3.74	3.34	3.11	2.96	2.85	2.70	2.53	2.44	2.39	2.31	2.27	2.24	2.19	2.13
15	4.54	3.68	3.29	3.06	2.90	2.79	2.64	2.48	2.39	2.33	2.25	2.21	2.18	2.12	2.07
16	4.49	3.63	3.24	3.01	2.85	2.74	2.59	2.42	2.33	2.28	2.20	2.16	2.13	2.07	2.01
17	4.45	3.59	3.20	2.96	2.81	2.70	2.55	2.38	2.29	2.23	2.15	2.11	2.08	2.02	1.96
18	4.41	3.55	3.16	2.93	2.77	2.66	2.51	2.34	2.25	2.19	2.11	2.07	2.04	1.98	1.92
19	4.38	3.52	3.13	2.90	2.74	2.63	2.48	2.31	2.21	2.15	2.07	2.02	2.00	1.94	1.88
20	4.35	3.49	3.10	2.87	2.71	2.60	2.45	2.28	2.18	2.12	2.04	1.99	1.96	1.90	1.84
22	4.30	3.44	3.05	2.82	2.66	2.55	2.40	2.23	2.13	2.07	1.98	1.93	1.91	1.84	1.78
24	4.26	3.40	3.01	2.78	2.62	2.51	2.36	2.18	2.09	2.03	1.94	1.89	1.86	1.80	1.73
26	4.23	3.37	2.98	2.74	2.59	2.47	2.32	2.15	2.05	1.99	1.90	1.85	1.82	1.76	1.69
28	4.20	3.34	2.95	2.71	2.56	2.45	2.29	2.12	2.02	1.96	1.87	1.81	1.78	1.72	1.65
30	4.17	3.32	2.92	2.69	2.53	2.42	2.27	2.09	1.99	1.93	1.84	1.79	1.76	1.69	1.62
40	4.08	3.23	2.84	2.61	2.45	2.34	2.18	2.00	1.90	1.84	1.74	1.69	1.66	1.59	1.51
50	4.03	3.18	2.79	2.56	2.40	2.29	2.13	1.95	1.85	1.78	1.69	1.63	1.60	1.52	1.44
60	4.00	3.15	2.76	2.53	2.37	2.25	2.10	1.92	1.81	1.75	1.65	1.59	1.56	1.48	1.39
70	3.98	3.13	2.74	2.50	2.35	2.23	2.07	1.89	1.79	1.72	1.62	1.56	1.53	1.45	1.35
80	3.96	3.11	2.72	2.48	2.33	2.21	2.05	1.88	1.77	1.70	1.60	1.54	1.51	1.42	1.32
100	3.94	3.09	2.70	2.46	2.30	2.19	2.03	1.85	1.75	1.68	1.57	1.51	1.48	1.39	1.28
150	3.91	3.06	2.67	2.43	2.27	2.16	2.00	1.82	1.71	1.64	1.54	1.47	1.44	1.34	1.22
200	3.89	3.04	2.65	2.41	2.26	2.14	1.98	1.80	1.69	1.62	1.52	1.45	1.42	1.32	1.19
400	3.86	3.02	2.62	2.39	2.23	2.12	1.96	1.78	1.67	1.60	1.49	1.42	1.38	1.28	1.13
200	3.84	2.99	2.60	2.37	2.21	2.09	1.94	1.75	1.64	1.57	1.46	1.40	1.32	1.24	1.00
	T.7.														

APPENDIX B

B-1 Analytical Methods for KOOl Regulated Organic Constituents

1711g/p.33

Regulated Constituent	Extraction Method	Method Number	Analytical Method/Met	hod No.
Volatile Organics				
To luene	Purge and trap	5030	GC/MS	82 40
Xylenes (total)	Purge and trap	5030	GC/MS	82 40
Semivolatile Organics				
Naphthalene	Continuous Liquid/ Extraction	3580	GC/MS	8270
Pentachlorophenol	Continuous Liquid/ Extraction	3580	GC/MS	8270
Phenanthrene	Continuous Liquid/ Extraction	3580	GC/MS	8270
Pyrene	Continuous Liquid/ Extraction	3580	GC/MS	8270

References: U.S. EPA. Test Methods for Evaluating Solid Waste, SW-846 Third Edition, Office of Solid Waste and Emergency Response, Washington, D.C. November, 1986.

Table B-2. K001 - Matrix Spike Recoveries Used to Calculate

Correction Factors for Regulated Organic Constituents

	Percent Recovery			
Regulated Constituent	Sample	Duplicate	Correction Factor Used	
Volatile Organics				
Toluene (ash)	99	110	1.01	
Toluene (water)	103	99	1.01	
Xylenes (ash)	86.4 (average)	96.4 (average)	1.16	
Xylenes (water)	100 (average)	94 (average)	1.15	
Semivolatile Organics				
Naphthalene (ash)	88 (average)	93 (avera ge)	1.14	
Naphthalene (water)	94 (average)	96 (avera ge)	1.06	
Pentachlorophenol (ash)	95	105	1.05	
Pentachlorophenol (water)	80	0	1.25	
Phenanthrene (ash)	88 (average)	93 (average)	1 14	
Phenanthrene (water)	94 (average)	96 (average)	1.06	
Pyrene (ash)	96	100	1.04	
Pyrene (water)	110	110	1.00	
Pyrene (water)	110	110	1.00	

Reference: Onsite Engineering Report for K001-Creosote

Table B-3 K001 Creosote Water Sample Volatile Organics Matrix Spike Recoveries (%)

	Spike level (ug/l)	#1	#2	RPD
1,1-Dichloroethene	25	100	97	3.0
Toluene	25	103	99	4.0
Chlorobenzene	25	95	90	5.4
Benzene	25	90	85	5 7
Trichloroethylene	25	69	64	7.5

Table B-4 K001 Creosote Ash Sample Volatile Organics Matrix Spike Recoveries (%)

	Spike level (ug/l)	#1	#2	RPD
1,1-Dichloroethene	25	86	95	9.9
To luene	25	99	110	10
Chlorobenzene	25	102	112	9.3
Benzene	25	78	88	12
Trichloroethylene	25	67	77	14

Table B-5 K001 Creosote Water Sample Semivolatile Organic Matrix Spike Recoveries (%)

Compound	Initial conc.	Amount added (ug/l)	% Re c	overy #2	RPD
Phenol	<10	100	65	61	6
2-Chlorophenol	<10	100	61	65	6
1,4-Dichlorobenzene	<10	50	61	51	16
N-nitroso-di-n-propylamine	<100	50	70	56	20
1,2,4-Trichlorobenzene	<10	50	72	54	25
4-Chloro-3-methylphenol	<10	100	80	73	9
Acenaphthene	<10	50	81	66	19
4-Nitrophenol	<50	100	0	0	NC
2,4-Dinitrotoluene	<10	50	21	17	19
Pentachlorophenol	<50	100	80	85	6
Pyrene	<10	50	62	60	3

NC - Not calculated.

Table B-6 K001 Creosote Ash Sample Semivolatile Organics Matrix Spike Recoveries (%)

	Initial conc.	Amount added	% Recovery		
Compound	(ug/g)	(ug/g)	#1	#2	RPD
Pheno 1	<4	67	72	65	9.7
2-Chlorophenol	<4	67	59	53	10
l,4-Dichlorobenzene	<4	33	48	46	4.2
N-nitroso-di-n-propylamine	<4	33	67	62	7.5
1,2,4-Trichlorobenzene	<4	33	30	30	0
4-Chloro-3-methylphenol	<4	67	35	€8	48
Acenaphthene	<4	33	0	3.4	100
4-Nitrophenol	<20	67	1.2	2.6	54
2,4-Dinitrotoluene	<4	33	0	0	NC
Pentachlorophenol	<20	67	0	0	NC
Pyrene	<4	33	0	0	NC

NC - Not calculated.

Table B-7 K001-PCP Ash Duplicate Matrix Spike Data, Volatile Organic Analyses

	Original amount	Amount	Amoun recovere <u>mg/lli</u>			
Compound	present, μg/liter	spiked, μg/liter	No 1	No 2	No 1	No. 2
Toluene	3	25	30	30	108	108
Chlorobenzene	<2	25	31	30	124	120
Benzene	<2	25	22	22	88	88
Trichloroethene	<2	25	22	21	88	84

Table B-8 K001-PCP TCLP Ash Duplicate Matrix Spike Data, Volatile Organic Analyses

	Original amount	Amount	Amou recove mg/l_l	red,	% Recovery	
Compound	present, μg/liter	spiked, μg/liter	No. 1	No. 2	No 1	No. 2
,1-Dichloraethylene	<2	25	23	28	92	112
oluene	<2	25	30	30	120	120
hlorobenzene	<2	25	30	26	120	104
enzene	<2	25	23	30	92	120
richloroethene	<2	25	21	21	84	84

Table B-9 KOO1-PCP TCLP Ash Duplicate Matrix Spike Data, Volatile Organic Analyses

	Initial concen- Amount		Amou recove mg/l l	% Recovery		
Compound	tration, μg/liter	added, μg/liter	No. 1	No. 2	No. 1	No 2
heno l	2	10,000	10,000	10,000	100	100
-Chlorophenol	<2	10,000	10,000	11,000	100	110
,4-Dichlorobenzene	<2	5,000	5,500	5,500	110	110
-Nitrosodinipropyl- amine	<5	5,000	6,500	7,000	130	140
,2,4-Trichlorobenzene	<5	5,000	4,700	4,600	94	92
-Chloro-3-methylphenol	< 5	10,000	8,000	9,000	80	90
cenaphthene	14,000	5,000	20,000	20,000	120	120
-Nitrophenol	<10	10,000	8,400	7,500	84	75
,4-Dinitrotoluene	<50	5,000	6,000	6,000	120	120
entachlorophenol	<50	10,000	10,000	10,000	100	100
yrene	11,000	5,000	15,000	16,000	80	100

Table B-10 K001-PCP TCLP Ash Duplicate Matrix Spike Data, Volatile Organic Analyses

	Initial concen-	Amount	Amou recove _mg/l l	red,	% Recovery	
Compound	tration, μg/liter	added, μg/liter	No. 1	No. 2	No. 1	No. 2
Pheno 1	2	200	170	160	85	80
-Chlorophenol	<2	200	200	210	100	105
.,4-Dichlorobenzene	<2	100	94	94	94	94
M-Nitrosodinipropyl- amine	<5	100	81	82	81	82
,2,4-Trichlorobenzene	<5	100	95	100	95	100
-Chloro-3-methylphenol	<5	200	180	190	90	95
Acenaphthene	<5	100	120	120	120	120
1-Nitrophenol	<10	200	200	180	100	90
2,4-Dinitrotoluene	<50	100	120	120	120	120
Pentachlorophenol	<50	200	190	210	95	105
Pyrene	<2	100	96	100	96	100

Table B-11 K001-PCP Scrubber Water Duplicate Matrix Spike Data Semivolatile Organic Analyses

	Initial concen-	Amount added,	Amou recove mq/l_l	red,	% Recovery	
Compound	tration, μg/liter	added, μg/liter	No. 1	No. 2	No. 1	No. 2
heno l	<2	200	60	45	30	22
-Chlorophenol	<2	200	56	40	28	20
,4-Dichlorobenzene	<2	100	85	87	85	87
-Nitrosodinipropyl- amine	< 5	100	70	66	70	66
,2,4-Trichlorobenzene	<5	100	110	120	110	120
-Chloro-3-methylphenol	<5	200	58	44	29	22
cenapht hene	<5	100	110	110	110	110
-Nitrophenol	<10	200	3	1.5	1.5	0.8
,4-Dinitrotoluene	<50	100	79	84	79	84
entachlorophenol	<50	200	6	3 4	3	1.7
yrene	<2	100	110	110	110	110

Table B-12 Matrix Spike Recoveries for Stabilized F006 Nonwastewater Residuals

Constituent	Original amount found (ppm)	Duplicate (ррт)	% Error	Actual Spike	% Recovery	Accuracy correction factor*
Copper	0 2247	0 2211	0.81	4 8494	92.5	1.08
	0 1526	0.1462	2.14	4 9981	97 0	1.03
Lead	0 3226	0.3091	2.14	4 9619	92.9	1.08
	0.2142	0.2287	3.27	4.6930	89.4	1.12
Zinc	0.0133	0.0238	28.3	5 0910	101.4	0.99
	27.202	3.65	76.3	19 818	87 8	1.14

^{*}Accuracy correction factor = 100 ÷ percent recovery.

Reference: Memo to R Turner, U.S. EPA/H.W.E R.L from Jesse R. Conner, Chemical Waste Management dated January 20, 1988

Table B-13 Matrix Spike Recovery for Metals for the TCLP Extract for K061 for Horsehead Resource Development Co

			Sample set 4			Sample set duplicate 4		
BDAT constituent	Original sample (ug/l)	Spike added (ug/l)	Spike result (ug/l)	Percent recovery*	Spike result (ug/l)	Percent recovery*	Relative percent difference (RPD)**	
Copper	<4 0	125	107	86	104	83	4	
Lead	< 5 0	25	22	88	19	76	15	
Zinc	2,640	10,000	12,600	100	12,400	98	2	

^{*} Percent Recovery = [(Spike Result ~ Original Amount)/Spike Amount] x 100

 $\label{eq:Reference: Onsite Engineering Report for Horsehead Resource Development Co.}$

^{**}RPD = $[(S1-S2)/(S1+S2)/2)] \times 100$, where S1 is the larger of the two percent recovery.

APPENDIX C

Table C-1

Methodology used by Agency to select regulated constituents for KOOl Volatiles and Semivolatiles is based on WCAPs for rotary kiln incineration. Namely, the concentration, boiling point, and bond dissociation energies are the basis for selection of regulated constituents.

	Concentration	on Boiling Point (°C)				Bond Dissociation Energy (Kcal/mol)			
	<u>Volatiles</u> (ppb)			<u>Volatiles</u>			Volatiles		
1.	Xylenes	120-170	1	Xylenes	140	1	Xylenes	1,905	
2	Toluene	10-170	2	Ethylbenzene	136	2.	Ethylbenzene	1,905	
3.	Ethylbenzene	55-87	3	Toluene	111	3	Toluene	1,620	
4	Benzene	51-83	4	Benzene	80	4.	Benzene	1,335	
	<u>Semivolatiles</u> (ppm)			iling point		Bond Dissociation Energy			
				(°C)			(Kcal/mol)		
1.	Naphtha lene	26,000-43,000	1	Benzo(b/k)	480	1.	Benzo(b/k	4,140	
				fluoranthrene			fluoranthrene		
2.	Phenanthrene	28,000-42,000	2	Chrysene	448	2	Benzo(c)pyrene	4,030	
3.	Acenaphtha lene	13,000-21,000	3.	Benz(a)anthracene	435	3.	Chrysene	3,775	
4	Fluoranthene	13,000-21,000	4	Pyrene	404	4	Benz(a)anthracene	3,680	
5 .	Fluorene	8,200-18,000	5	Fluoranthene	375	5.	Pyrene	3,240	
6.	Pyrene	12,000-17,000	6	Anthrecene	342	6.	Fluoranthene	3,130	
7.	Anthracene	7,300-15,000	7	Phenanthrene	340	7.	Phenanthrene	2,900	
8	Chrysene	2,500-4,800	8	Benzo(a)pyrene	311	8.	Fluorene	2,725	
9	Phenol	2,400-3,900	9	Pentachlorophenol	309	9.	Anthracene	2,715	
10.	Benz(a)anthracene	2,500-3,400	10.	Fluorene	295	10	Acenaphene	2,570	
11.	Pentachlorophenol	920-3,000	11.	Acenaphthene	279	11	Naphthalene	2,025	
12	Benzo(b/k)		12.	Naphthalene	218	12	Pheno 1	1,435	
	fluoranthrene	940-2,300	13.	Pheno 1	182	13.	Pentachlorophenol	1,410	
13	Benzo(a)pyrene	250-940					•		

APPENDIX D

Analytical Method for Determining the Thermal Conductivity of a Waste

APPENDIX D

The comparative method of measuring thermal conductivity has been proposed as an ASTM test method under the name "Guarded, Comparative, Longitudinal Heat Flow Technique." A thermal heat flow circuit is used which is the analog of an electrical circuit with resistances in series. A reference material is chosen to have a thermal conductivity close to that estimated for the sample. Reference standards (also known as heat meters) having the same cross-sectional dimensions as the sample are placed above and below the sample. An upper heater, a lower heater, and a heat sink are added to the "stack" to complete the heat flow circuit. See Figure 1.

The temperature gradients (analogous to potential differences) along the stack are measured with type K (chromel/alumel) thermocouples placed at known separations. The thermocouples are placed into holes or grooves in the references and also in the sample whenever the sample is thick enough to accommodate them.

For molten samples, pastes, greases, and other materials that must be contained, the material is placed into a cell consisting of a top and bottom of Pyrex 7740 and a containment ring of marinite. The sample is 2 inch in diameter and .5 inch thick. Thermocouples are not placed into the sample but rather the temperatures measured in the Pyrex are extrapolated to give the temperature at the top and bottom surfaces of the sample material. The Pyrex disks also serve as the thermal conductivity reference material.

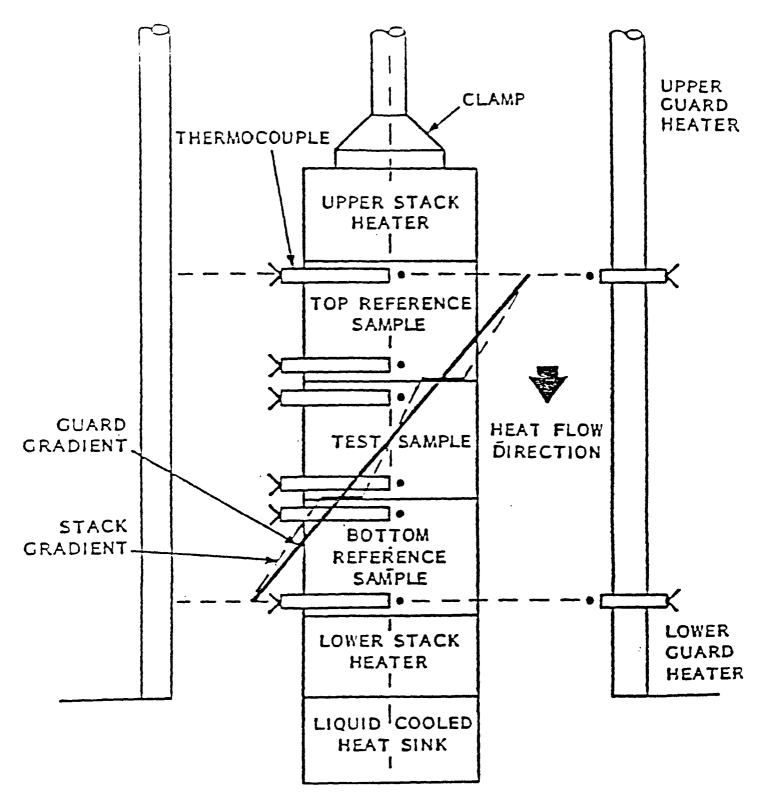


Figure D-1
SCHEMATIC DIAGRAM OF THE COMPARATIVE METHOD

The stack is clamped with a reproducible load to insure intimate contact between the components. In order to produce a linear flow of heat down the stack and reduce the amount of heat that flows radially, a guard tube is placed around the stack and the intervening space is filled with insulating grains or powder. The temperature gradient in the guard is matched to that in the stack to further reduce radial heat flow.

The comparative method is a steady state method of measuring thermal conductivity. When equilibrium is reached the heat flux (analogous to current flow) down the stack can be determined from the references. The heat into the sample is given by

$$Q_{in} = \lambda_{top} (dT/dx)_{top}$$

and the heat out of the sample is given by

$$\int_{0}^{Q} out = \lambda_{bottom} (dT/dx)_{bottom}$$

where

 λ = thermal conductivity

dT/dx = temperature gradient

and top refers to the upper reference while bottom refers to the lower reference. If the heat was confined to flow just down the stack, then Q_{in} and Q_{out} would be equal. If Q_{in} and Q_{out} are in reasonable agreement, the average heat flow is calculated from

$$Q = (Q_{in} + Q_{out})/2$$

The sample thermal conductivity is then found from

$$\lambda_{\text{sample}} = Q/(dT/dx)_{\text{sample}}$$

APPENDIX E CALCULATIONS OF TREATMENT STANDARDS



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1 Tolvere

Woter USL

$$<50$$

 <10 0.050 pm
99
 100 CF=1.01
 $CF=1.01$
 $CF=1.01$

2 Xylers

Detection Limits Ash HLO Use.

Crossole OFR 450 <50 <0.050 pp.

PCP OFR 410 <10

CF(ash = 1.16)

Recomp for Crossole = 86.4 87 CF(Ho = 1.15)

$$\mu$$
 PCP = 94 93.6 VF= 2.8

Xyleres Standard Ash = 0.162 H20 = 0.161

Standard Rounded Off to 2 Significant Figures

Tolvene (ash) 0.14 mg/kg
Tolvene (veter) 0.14 mg/kg

Xylenes (ash) 0.16 mg/kg

Xylenes (water) 0.16 mg/k

Wernaring.

6850 Versar Center Springfield, VA 22151 (703) 750-3000

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100	~	Sei	ni-Voas		PROJECT	 				JOB NO	DAT	E
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3 Nophtholese

Detection Limits	Ash	Water	Use
CreosofuER	< 0.55	<0.010	CUSS 8
PCP OER	12.5	< 0.050	< 2.5 ash
			< 0.050 water
Recogns for Naphllu			
Crosute	88	57.4	CF(ash)= 羅 1.17
PLP	88	94	CF (pcp) = \$ 1.06
			VF=2.8

Naphtholen Stanlar Ash = 7.98 " H20 = 0.148

(4) Phenowthren

Standards Rounded Off to 2 significant Figures

Naphthelere (ash) 8.0 ms/kg
Naphthelere (water) 0.15 ms/k
Phenonthrene (ash) 8.0 ms/ks
Phenonthrene (water) 0.15 ms/k

Phenather Starled Ash = 7.98 H10=0.148



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5 Pentachlorophenol

Detection Limits

Ash weter <12.5 <0.25 Use ASL < 125 HIU < 0.250

Recorney PCP = 95

woter 3, 1.7*

CF(aoL) = 1.05 CF(HO) = 1.25VF = 2.8

Pertachlorphal Stanland Ash=36.75

@ Pyrene

Detective Limits Creasok Ash Welco <0.65 <0.010 <2.5 <0.050

 $\frac{USC}{ASL}$ < 2.5 H_{20} < 0.050

Recovers for Pyrene Greasoft PCP

0 60 96 110 CF(ash) = 1.04 CF(H20)= 日本期 VF=2.8

Stoleto Rankel Off to 2 signified figure

PCP (cosh) 37 mg/kg PCP (water) 0.88 mg/l Pyrene (cosh) 7.3 mg/kg

Pyrere (water) 0.14 mg/l

Pyrene Stanbard Ash = 7.28 " H.O = 0.140



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Lead

Accuracy Corrected Values	Variability Factor	Student
0.39	1.46	0.45 mg/kg
0.23		-
0,29		
0.34		
0.34		
0.35		
0,34		
0.27		

mean = 0.32

Copper		
0.21	2.46	0.71 mg 1kg
0.15		
0.21		
0.07		
0.14		
0.12		
0.16		
0.16		
0.08		
0.14	•	
0.24		

Mean = 0.18



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ZINC

Accuracy Corrected Values Variability Factor Standard

0.03

1.46

5,45 mg/kg

0.01 0.01 0.03 0.02

0.03

0.02

mea = 0.02

8 i.



SUBJECT (1001 Waskwoters Metals dota from Envinte			CLIENT			BY	BY				
Metals	dc	ナC	m Envi	4 K	PROJECT				JOB NO	DAT	E
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According Corrected Victor	Variability Factor	Stanlard
0.25	2.31	0.42 mg/l
0.18		•
0.25		
0.03		
0.17		
0.14		
0.19		
0.19		
0.10		
0.17		
c. 29		
may - 0.18		

L

mcin = 0.18		
Leal		
<0.013	2.8	0.037 mg/1
< 0.013		
0.013		
0.013		
0.013		
0,013		
0,013		
0.013		
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metab Detafm Evink	PROJECT	JOB NO DATE
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Zinc

According Corrected Value

Variability Factor

4.13

Stadad

10

0.128

0.117

0.143

1.653

0.128

0.097

0.117

0.133

0.041

0.071

0.102

mcon = 0.250